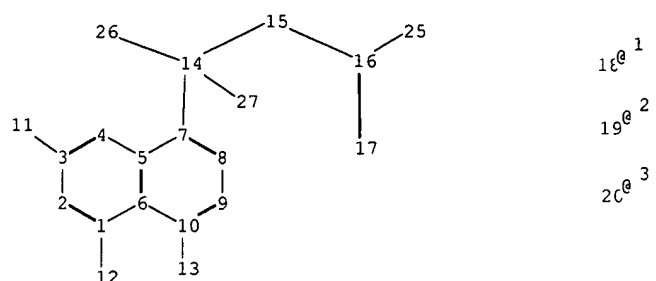


O @<sup>1</sup>  
N @<sup>2</sup>  
S @<sup>3</sup>

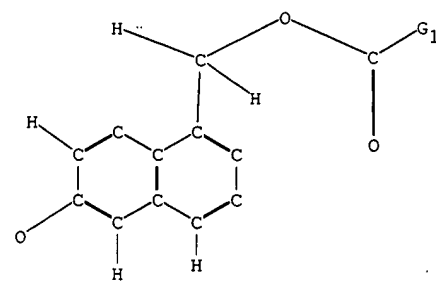


1@<sup>1</sup>  
19@<sup>2</sup>  
20@<sup>3</sup>

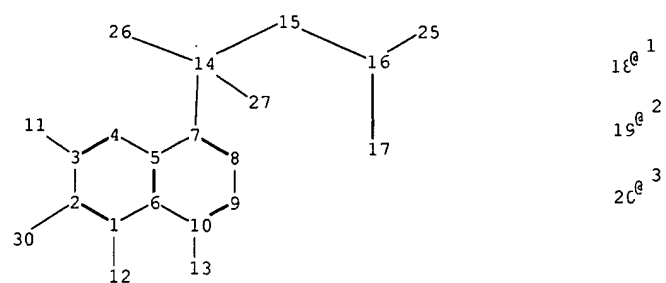
chain nodes :  
11 12 13 14 15 16 17 25 26 27  
ring nodes :  
1 2 3 4 5 6 7 8 9 10  
ring/chain nodes :  
18 19 20  
chain bonds :  
1-12 3-11 7-14 10-13 14-15 14-26 14-27 15-16 16-17 16-25  
ring bonds :  
1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-10 7-8 8-9 9-10  
exact/norm bonds :  
14-15 15-16 16-17 16-25  
exact bonds :  
1-12 3-11 7-14 10-13 14-26 14-27  
normalized bonds :  
1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-10 7-8 8-9 9-10

G1:[\*1],[\*2],[\*3]

Match level :  
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:CLASS  
12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS  
25:CLASS 26:CLASS 27:CLASS



O @<sup>1</sup>  
N @<sup>2</sup>  
S @<sup>3</sup>



1E@<sup>1</sup>  
19@<sup>2</sup>  
2C@<sup>3</sup>

chain nodes :  
11 12 13 14 15 16 17 25 26 27 30  
ring nodes :  
1 2 3 4 5 6 7 8 9 10  
ring/chain nodes :  
18 19 20  
chain bonds :  
1-12 2-30 3-11 7-14 10-13 14-15 14-26 14-27 15-16 16-17 16-25  
ring bonds :  
1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-10 7-8 8-9 9-10  
exact/norm bonds :  
2-30 14-15 15-16 16-17 16-25  
exact bonds :  
1-12 3-11 7-14 10-13 14-26 14-27  
normalized bonds :  
1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-10 7-8 8-9 9-10  
isolated ring systems :  
containing 1 :

G1:[\*1],[\*2],[\*3]

Match level :  
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:CLASS  
12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS  
25:CLASS 26:CLASS 27:CLASS 30:CLASS

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PASSWORD:  
TERMINAL (ENTER 1, 2, 3, OR ?):2
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\* \* \* \* \* \* \* \* \* \* Welcome to STN International \* \* \* \* \* \* \* \* \* \*

NEWS 1 Web Page URLs for STN Seminar Schedule - N. America  
NEWS 2 Dec 17 The CA Lexicon available in the CAPLUS and CA files  
NEWS 3 Feb 06 Engineering Information Encompass files have new names  
NEWS 4 Feb 16 TOXLINE no longer being updated  
NEWS 5 Apr 23 Search Derwent WPINDEX by chemical structure  
NEWS 6 Apr 23 PRE-1967 REFERENCES NOW SEARCHABLE IN CAPLUS AND CA  
NEWS 7 May 07 DGENE Reload  
NEWS 8 Jun 20 Published patent applications (A1) are now in USPATFULL  
NEWS 9 JUL 13 New SDI alert frequency now available in Derwent's  
DWPI and DPCI

NEWS EXPRESS July 11 CURRENT WINDOWS VERSION IS V6.0b,  
CURRENT MACINTOSH VERSION IS V5.0C (ENG) AND V5.0JB (JP),  
AND CURRENT DISCOVER FILE IS DATED 06 APRIL 2001

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|---------------------|------------------|
| 0.15                | 0.15             |

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DICTIONARY FILE UPDATES: 2 AUG 2001 HIGHEST RN 350221-07-7

TSCA INFORMATION NOW CURRENT THROUGH January 11, 2001

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Structure search limits have been increased. See HELP SLIMIT for details.

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Uploading 09633697.str

L1        STRUCTURE UPLOADED

=> s 11

SAMPLE SEARCH INITIATED 10:54:12 FILE 'REGISTRY'  
SAMPLE SCREEN SEARCH COMPLETED - 246 TO ITERATE

100.0% PROCESSED      246 ITERATIONS  
SEARCH TIME: 00.00.02

11 ANSWERS

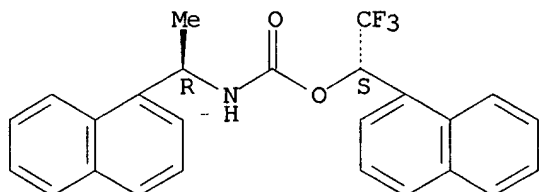
FULL FILE PROJECTIONS:    ONLINE    \*\*COMPLETE\*\*  
                                  BATCH    \*\*COMPLETE\*\*  
PROJECTED ITERATIONS:      3979 TO      5861  
PROJECTED ANSWERS:          21 TO      417

L2        11 SEA SSS SAM L1

=> d scan

L2    11 ANSWERS    REGISTRY    COPYRIGHT 2001 ACS  
IN    Carbamic acid, [1-(1-naphthalenyl)ethyl]-, 2,2,2-trifluoro-1-(1-naphthalenyl)ethyl ester, [S-(R\*,S\*)]- (9CI)  
MF    C25 H20 F3 N O2

Absolute stereochemistry.



HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):

Uploading

'UPLOAD SSTN' IS NOT VALID HERE

To display more answers, enter the number of answers you would like to see. To end the display, enter "NONE", "N", "0", or "END".

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):09633697.str

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):

'0 SZ' @-#&1~" J\* ' IS NOT VALID HERE

To display more answers, enter the number of answers you would like to see. To end the display, enter "NONE", "N", "0", or "END".  
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):  
'0 SZ' @-#&1~" J\* ' IS NOT VALID HERE

To display more answers, enter the number of answers you would like to see. To end the display, enter "NONE", "N", "0", or "END".  
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):  
'0 SZ' @-#&1~" J\* ' IS NOT VALID HERE

To display more answers, enter the number of answers you would like to see. To end the display, enter "NONE", "N", "0", or "END".  
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):  
'0 SZ' @-#&1~" J\* ' IS NOT VALID HERE

To display more answers, enter the number of answers you would like to see. To end the display, enter "NONE", "N", "0", or "END".  
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=>  
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### L3 STRUCTURE UPLOADED

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SAMPLE SCREEN SEARCH COMPLETED - 246 TO ITERATE

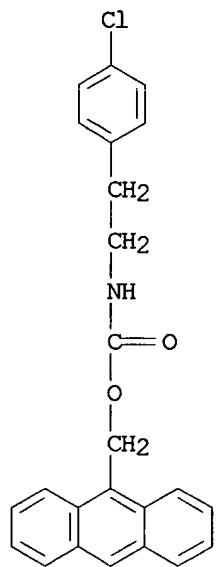
100.0% PROCESSED 246 ITERATIONS 5 ANSWERS  
SEARCH TIME: 00.00.03

|                        |         |              |
|------------------------|---------|--------------|
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|                        | BATCH   | **COMPLETE** |
| PROJECTED ITERATIONS:  | 3979 TO | 5861         |
| PROJECTED ANSWERS:     | 5 TO    | 234          |

L4 5 SEA SSS SAM L3

=> d scan

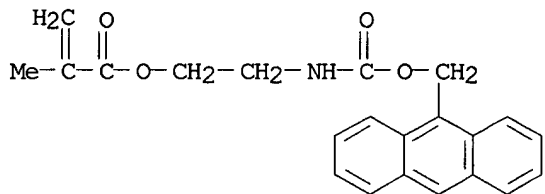
L4 5 ANSWERS REGISTRY COPYRIGHT 2001 ACS  
IN Carbamic acid, [2-(4-chlorophenyl)ethyl]-, 9-anthracenylmethyl ester  
(9CI)  
MF C24 H20 Cl N O2



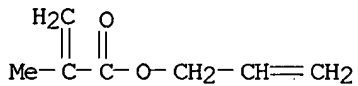
HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L4 5 ANSWERS REGISTRY COPYRIGHT 2001 ACS  
 IN 2-Propenoic acid, 2-methyl-,  
 2-[[(9-anthracenylmethoxy)carbonyl]amino]ethyl ester, polymer with 2-propenyl 2-methyl-2-propenoate (9CI)  
 MF (C22 H21 N O4 . C7 H10 O2)x  
 CI PMS

CM 1



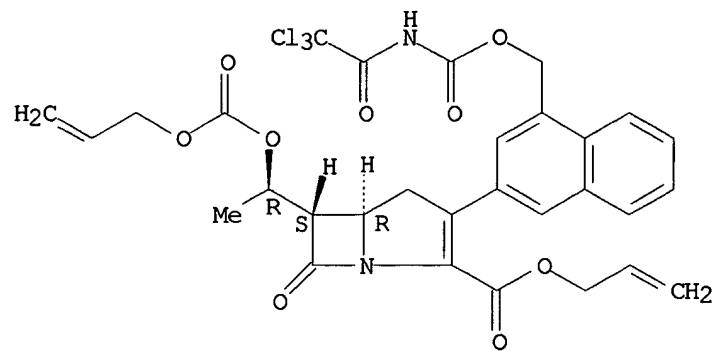
CM 2



L4 5 ANSWERS REGISTRY COPYRIGHT 2001 ACS  
 IN 1-Azabicyclo[3.2.0]hept-2-ene-2-carboxylic acid, 7-oxo-6-[1-[[[2-  
 propenyloxy)carbonyl]oxy]ethyl]-3-[4-[[[[trichloroacetyl]amino]carbonyl]o

xy]methyl]-2-naphthalenyl]-, 2-propenyl ester, [5R-[5.alpha.,6.alpha.(R\*)]]- (9CI)  
MF C30 H27 Cl3 N2 O9

Absolute stereochemistry.



HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

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L5 STRUCTURE UPLOADED

=> s 15

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SAMPLE SCREEN SEARCH COMPLETED - 246 TO ITERATE

100.0% PROCESSED 246 ITERATIONS  
SEARCH TIME: 00.00.01

3 ANSWERS

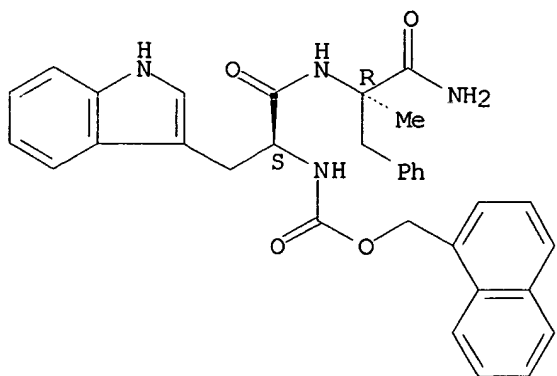
FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*  
BATCH \*\*COMPLETE\*\*  
PROJECTED ITERATIONS: 3979 TO 5861  
PROJECTED ANSWERS: 3 TO 162

L6 3 SEA SSS SAM L5

=> d scan

L6 3 ANSWERS REGISTRY COPYRIGHT 2001 ACS  
IN D-Phenylalaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-  
.alpha.-methyl- (9CI)  
MF C33 H32 N4 O4

Absolute stereochemistry.

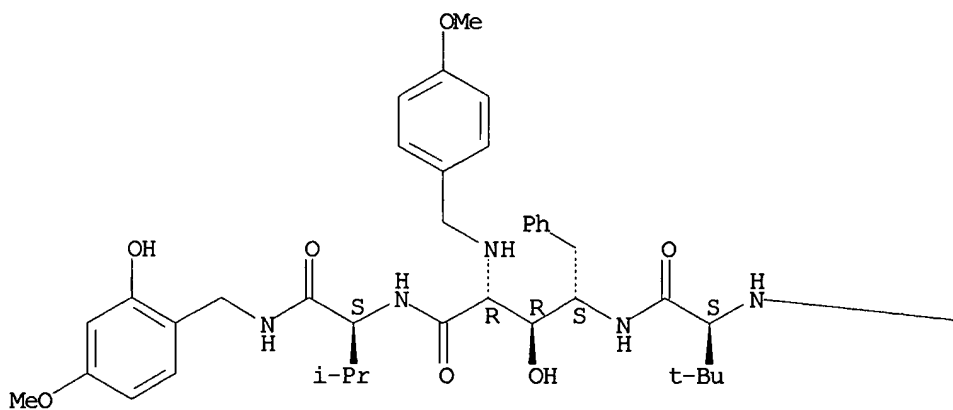


HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):2

L6 3 ANSWERS REGISTRY COPYRIGHT 2001 ACS  
 IN L-Valinamide,  
 N4-[3-methyl-N-[(1-naphthalenylmethoxy)carbonyl]-L-valyl]-4-  
 amino-2,4,5-trideoxy-2-[(4-methoxyphenyl)methyl]amino]-5-phenyl-L-  
 lyxonoyl-N-[(2-hydroxy-4-methoxyphenyl)methyl]- (9CI)  
 MF C50 H61 N5 O9

Absolute stereochemistry.

PAGE 1-A

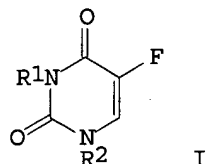


L2 ANSWER 1 OF 1 CA COPYRIGHT 2001 ACS

102:12247 Prodrugs of 5-fluorouracil. II. Hydrolysis kinetics, bioactivation, solubility and lipophilicity on N-alkoxycarbonyl derivatives of 5-fluorouracil. Buur, Anders; Bundgaard, Hans (Dep. Pharm.

Chem., R. Dan. Sch. Pharm., Copenhagen, DK-2100, Den.). Arch. Pharm. Chemi, Sci. Ed., 12(2), 37-44 (English) 1984. CODEN: AVPCCS. ISSN: 0302-248X.

GI



AB The decompn. and bioactivation characteristics of 5 N3- and N1,N3-alkoxycarbonyl derivs. (I, R1 = CO2Ph, CO2CH2Ph, or CO2Et and R2 = H, CO2Ph, or CO2CH2Ph) of 5-fluorouracil [51-21-8] were studied to assess

their suitability as prodrugs for the parent compd. The N1,N3-disubstituted derivs. were very unstable in aq. soln. and were subject to spontaneous and hydroxide ion-catalyzed hydrolysis with formation of the corresponding N3-deriv. The half-life for the selective removal of the N1-alkoxycarbonyl group was 2 min at pH 1-7 and 37 .degree.. The N3-alkoxycarbonyl group was highly resistant towards chem. hydrolysis, but showed enzyme-mediated cleavage in human plasma and, in particular, rat liver homogenate. The N3-alkoxycarbonyl derivs. were more

lipophilic than 5-fluorouracil as detd. by partition expts. in octanol-aq. buffer systems but as shown for the N3-ethoxycarbonyl deriv., the aq. solv. was at the same time greatly enhanced. Thus, N3-alkoxycarbonyl derivs. may be considered as potentially useful prodrug forms of 5-fluorouracil, although it may be questioned whether their conversion to the parent drug is sufficiently facile under in vivo conditions.

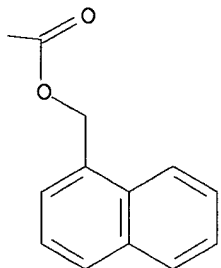
L3 ANSWER 1 OF 1 CA COPYRIGHT 2001 ACS

119:62362 Chemical and biological degradation of 5-fluorouracil prodrugs having high serum albumin binding potencies. Suda, Yasuo; Shimidzu, Kenji; Sumi, Masao; Kusumoto, Shoichi; Nadai, Tanekazu; Yamashita, Shinji (Fac. Sci., Osaka Univ., Toyonaka, 560, Japan). Biol. Pharm. Bull., 16(3), 322-4 (English) 1993. CODEN: BPBLEO.

AB In order to understand the fundamental structural features which yield both high serum albumin binding potency and desired property as a prodrug,

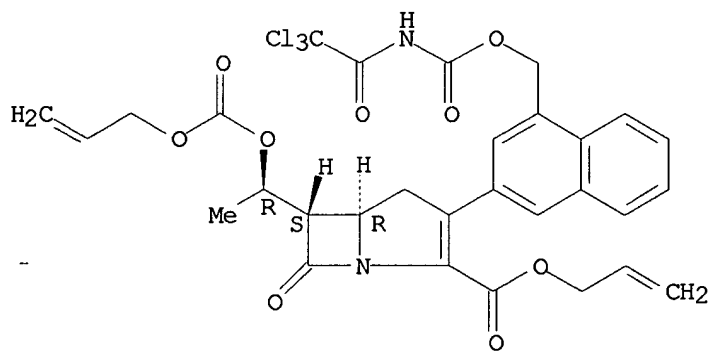
the derivatization was performed at N-1 or N-3 position in 5-fluorouracil.

The N-3 derivs. were more stable than N-1 derivs. in vitro, whereas they were metabolized quickly in vivo. It is suggested that N-1 position should be blocked to avoid fast metab. in vivo.



L6 3 ANSWERS REGISTRY COPYRIGHT 2001 ACS  
IN 1-Azabicyclo[3.2.0]hept-2-ene-2-carboxylic acid, 7-oxo-6-[1-[(2-  
propenoxy)carbonyl]oxyethyl]-3-[4-[[[[trichloroacetyl]amino]carbonyl]o  
xy]methyl]-2-naphthalenyl-, 2-propenyl ester, [5R-  
[5.alpha.,6.alpha.(R\*)]]- (9CI)  
MF C<sub>30</sub> H<sub>27</sub> Cl<sub>3</sub> N<sub>2</sub> O<sub>9</sub>

Absolute stereochemistry.



ALL ANSWERS HAVE BEEN SCANNED

=> d his

(FILE 'HOME' ENTERED AT 10:52:11 ON 03 AUG 2001)

FILE 'REGISTRY' ENTERED AT 10:52:21 ON 03 AUG 2001

L1 STRUCTURE uploaded  
L2 11 S L1

L3                   STRUCTURE UPLOADED  
L4                 5 S L3  
L5                   STRUCTURE UPLOADED  
L6                 3 S L5

=> s 15 sss full  
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FULL SCREEN SEARCH COMPLETED - 5189 TO ITERATE

100.0% PROCESSED   5189 ITERATIONS  
SEARCH TIME: 00.00.02

73 ANSWERS

L7                 73 SEA SSS FUL L5

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L8                   STRUCTURE UPLOADED

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FULL SUBSET SCREEN SEARCH COMPLETED - 13 TO ITERATE

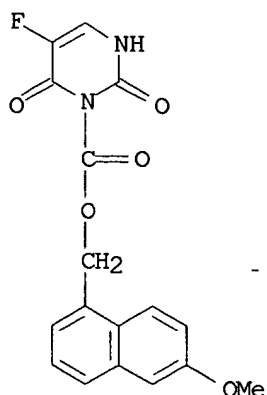
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SEARCH TIME: 00.00.01

3 ANSWERS

L9                 3 SEA SUB=L7 SSS FUL L8

=> d 1-3 ide cbib

L9   ANSWER 1 OF 3 REGISTRY COPYRIGHT 2001 ACS  
RN   238761-23-4 REGISTRY  
CN   1(2H)-Pyrimidinecarboxylic acid, 5-fluoro-3,6-dihydro-2,6-dioxo-,  
      (6-methoxy-1-naphthalenyl)methyl ester (9CI) (CA INDEX NAME)  
OTHER NAMES:  
CN   DMU 339  
FS   3D CONCORD  
MF   C17 H13 F N2 O5  
SR   CA  
LC   STN Files: CA, CAPLUS, TOXLIT



1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 131:175072 Hydroxylation-activated drug release, and prodrug preparation. Potter, Gerard Andrew; Patterson, Lawrence Hylton; Burke, Michael Danny (De Montfort University, UK). PCT Int. Appl. WO 9940944 A2 19990819, 53 pp. DESIGNATED STATES: W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG. (English). CODEN: PIXXD2. APPLICATION: WO 1999-GB416 19990210. PRIORITY: GB 1998-2957 19980212; US 1998-115016 19980714.

L9 ANSWER 2 OF 3 REGISTRY COPYRIGHT 2001 ACS

RN 238761-22-3 REGISTRY

CN Carbamic acid, bis(2-chloroethyl)-, (6-methoxy-1-naphthalenyl)methyl ester

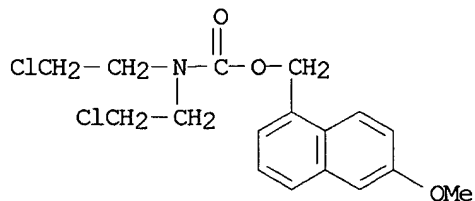
(9CI) (CA INDEX NAME)

FS 3D CONCORD

MF C17 H19 Cl2 N O3

SR CA

LC STN Files: CA, CAPLUS, TOXLIT



1 REFERENCES IN FILE CA (1967 TO DATE)

1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 131:175072 Hydroxylation-activated drug release, and prodrug preparation. Potter, Gerard Andrew; Patterson, Lawrence Hylton; Burke, Michael Danny (De Montfort University, UK). PCT Int. Appl. WO 9940944 A2 19990819, 53 pp. DESIGNATED STATES: W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG. (English). CODEN: PIXXD2. APPLICATION: WO 1999-GB416 19990210. PRIORITY: GB 1998-2957 19980212; US 1998-115016 19980714.

L9 ANSWER 3 OF 3 REGISTRY COPYRIGHT 2001 ACS

RN 238761-21-2 REGISTRY

CN Carbamic acid, [(7S)-5,6,7,9-tetrahydro-1,2,3,10-tetramethoxy-9-oxobenzo[a]heptalen-7-yl]-, (6-methoxy-1-naphthalenyl)methyl ester (9CI) (CA INDEX NAME)

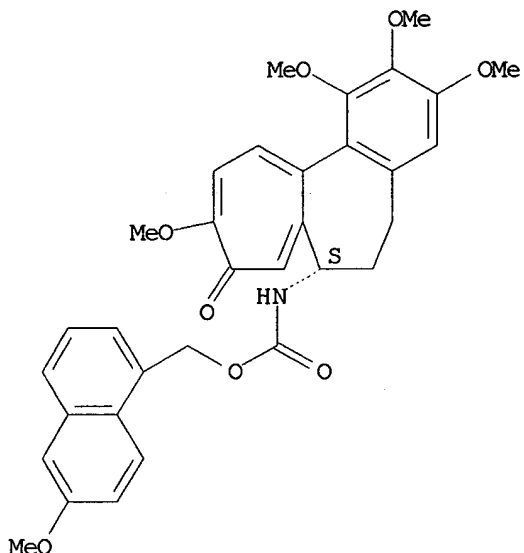
OTHER NAMES:

CN DMU 331

FS STEREOSEARCH

MF C33 H33 N O8  
SR CA  
LC STN Files: CA, CAPLUS, TOXLIT

Absolute stereochemistry.



1 REFERENCES IN FILE CA (1967 TO DATE)  
1 REFERENCES IN FILE CAPLUS (1967 TO DATE)

REFERENCE 1: 131:175072 Hydroxylation-activated drug release, and prodrug preparation. Potter, Gerard Andrew; Patterson, Lawrence Hylton; Burke, Michael Danny (De Montfort University, UK). PCT Int. Appl. WO 9940944 A2 19990819, 53 pp. DESIGNATED STATES: W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG. (English). CODEN: PIXXD2. APPLICATION: WO 1999-GB416 19990210. PRIORITY: GB 1998-2957 19980212; US 1998-115016 19980714.

=> s 17 not 19  
L10 70 L7 NOT L9

=> file caplus  
COST IN U.S. DOLLARS  
FULL ESTIMATED COST

| SINCE FILE ENTRY | TOTAL SESSION |
|------------------|---------------|
| 176.17           | 176.32        |

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FILE LAST UPDATED: 2 Aug 2001 (20010802/ED)

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L11      50 L10

=> s l11 and p/dt
      3240976 P/DT
L12      32 L11 AND P/DT

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      8723098 ED<19981202
      (ED<981202)
L13      11 L11 NOT P/DT AND ED<19981202

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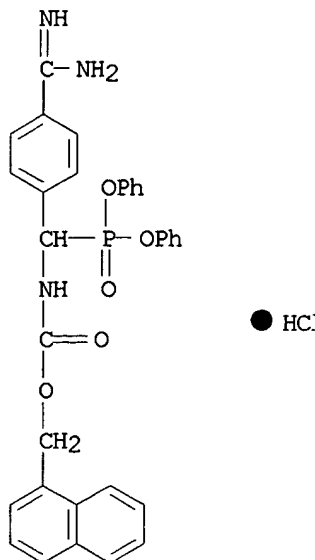
L13 ANSWER 1 OF 11 CAPLUS COPYRIGHT 2001 ACS
1998:338712 Document No. 129:95705 Synthesis and Evaluation of Diphenyl
      Phosphonate Esters as Inhibitors of the Trypsin-like Granzymes A and K
and
      Mast Cell Tryptase. Jackson, Delwin S.; Fraser, Stephanie A.; Ni,
      Li-Ming; Kam, Chih-Min; Winkler, Ulrike; Johnson, David A.; Froelich,
      Christopher J.; Hudig, Dorothy; Powers, James C. (School of Chemistry and
      Biochemistry, Georgia Institute of Technology, Atlanta, GA, 30332-0400,
      USA). J. Med. Chem., 41(13), 2289-2301 (English) 1998. CODEN: JMCMAR.
      ISSN: 0022-2623. Publisher: American Chemical Society.
IT 209675-92-3P
      RL: BAC (Biological activity or effector, except adverse); RCT
      (Reactant);
      SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
```

(prepn. and structure-activity of phosphonate ester inhibitors of the trypsin-like granzymes A and K and mast cell tryptase)

RN 209675-92-3 CAPLUS

CN Carbamic acid,

[ [4-(aminoiminomethyl)phenyl] (diphenoxypyrophosphinyl)methyl]-, 1-naphthalenylmethyl ester, monohydrochloride (9CI) (CA INDEX NAME)

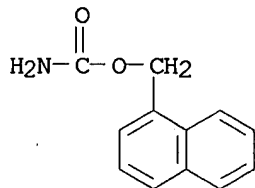


IT 74156-18-6P 209675-90-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)  
(prepn. and structure-activity of phosphonate ester inhibitors of the trypsin-like granzymes A and K and mast cell tryptase)

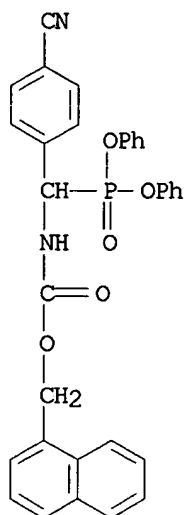
RN 74156-18-6 CAPLUS

CN 1-Naphthalenemethanol, carbamate (9CI) (CA INDEX NAME)



RN 209675-90-1 CAPLUS

CN Carbamic acid, [(4-cyanophenyl) (diphenoxypyrophosphinyl)methyl]-, 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)



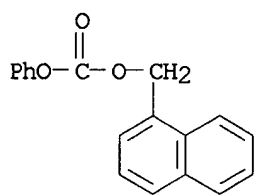
L13 ANSWER 2 OF 11 CAPLUS COPYRIGHT 2001 ACS  
 1994:667467 Document No. 121:267467 The photochemistry of 1-naphthylmethyl carbonates and carbamates. Parman, T.; Pincock, J. A.; Wedge, P. J.  
 (Dep.

Chem., Dalhousie, Halifax, NS, B3H 4J3, Can.). Can. J. Chem., 72(5), 1254-61 (English) 1994. CODEN: CJCHAG. ISSN: 0008-4042.

IT 158833-25-1P 158833-26-2P  
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)  
 (excited state properties and photochem. of)

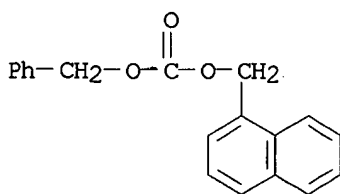
RN 158833-25-1 CAPLUS

CN Carbonic acid, 1-naphthylmethyl phenyl ester (9CI) (CA INDEX NAME)



RN 158833-26-2 CAPLUS

CN Carbonic acid, 1-naphthylmethyl phenylmethyl ester (9CI) (CA INDEX NAME)



L13 ANSWER 3 OF 11 CAPLUS COPYRIGHT 2001 ACS

1994:631319 Document No. 121:231319 Rational design of high affinity tachykinin NK2 receptor antagonists. Boyle, S.; Guard, S.; Hodgson, J.; Horwell, D. C.; Howson, W.; Hughes, J.; McKnight, A.; Martin, K.; Pritchard, M. C.; et al. (Parke-Davis Neurosci. Res. Cent., Addenbrookes Hosp. Site, Cambridge, CB2 2QB, UK). Bioorg. Med. Chem., 2(2), 101-13 (English) 1994. CODEN: BMECEP. ISSN: 0968-0896.

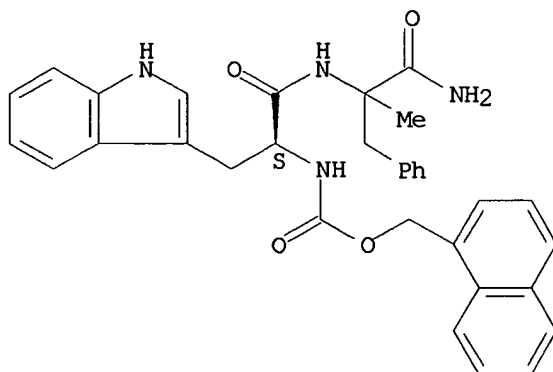
IT **146034-77-7P**

RL: SPN (Synthetic preparation); PREP (Preparation)  
(prepn. and neurokinin-2 receptor binding affinity of)

RN 146034-77-7 CAPLUS

CN Phenylalaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L13 ANSWER 4 OF 11 CAPLUS COPYRIGHT 2001 ACS

1994:243889 Document No. 120:243889 Synthesis and study of the properties of

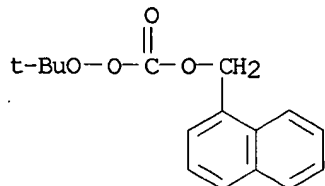
a new series of OO-tert-butyl O-(alkylbenzyl) and O-(naphthylmethyl) peroxy carbonates. Etlis, I. V.; Fomin, V. A.; Nozrina, F. D.; Kurskii, Yu. A.; Shmuilovich, S. M. (Gos. Nauchno-Issled. Inst. Khim. Tekhnol. Polimer., Dzerzhinsk, Russia). Zh. Org. Khim., 29(5), 994-1000 (Russian) 1993. CODEN: ZORKAE. ISSN: 0514-7492.

IT **154422-60-3P**

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)  
(prepn. and thermolysis kinetics of)

RN 154422-60-3 CAPLUS

CN Carbonoperoxoic acid, OO-(1,1-dimethylethyl) O-(1-naphthalenylmethyl) ester (9CI) (CA INDEX NAME)



L13 ANSWER 5 OF 11 CAPLUS COPYRIGHT 2001 ACS

1994:216500 Document No. 120:216500 Study of the thermal decomposition of bis(alkylbenzyl) and bis(naphthylmethyl) peroxydicarbonates as a function of the structure of the alkylaromatic fragments. Fomin, V. A.; Etlis, I. V.; Kurskii, Yu. A.; Nozrina, F. D.; Chervyakova, G. N.; Shmuilovich, S. M. (NII Khim Tekhnol. Polim., Dzerzhinsk, Russia). Zh. Org. Khim.,

29(5),

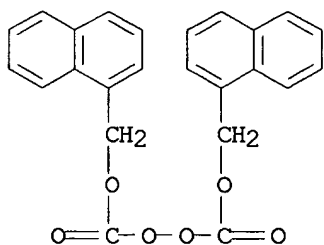
982-93 (Russian) 1993. CODEN: ZORKAE. ISSN: 0514-7492.

IT 138556-70-4 138556-73-7

RL: PRP (Properties); RCT (Reactant)  
(thermal decomprn. of, kinetics of)

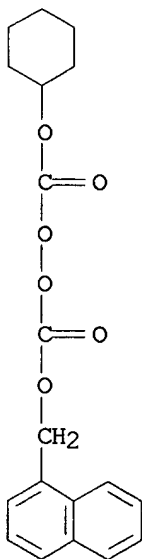
RN 138556-70-4 CAPLUS

CN Peroxydicarbonic acid, bis(1-naphthalenylmethyl) ester (9CI) (CA INDEX NAME)



RN 138556-73-7 CAPLUS

CN Peroxydicarbonic acid, cyclohexyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)



L13 ANSWER 6 OF 11 CAPLUS COPYRIGHT 2001 ACS

1993:234530 Document No. 118:234530 Process controlling of vinyl chloride polymerization in mass (suspension) with high degree of conversion.

Grishin, A. N.; Zegelman, V. I.; Fomin, V. A.; Etlis, I. V.; Popov, V. A.;

Khavritsyn, A. A. (Res. Inst. Polym. Chem. Technol., Dzerzhinsk, Russia).  
DECHEMA Monogr., 127(Int. Workshop Polym. React. Eng., 4th, 1992), 449-59  
(English) 1992. CODEN: DMDGAG. ISSN: 0070-315X.

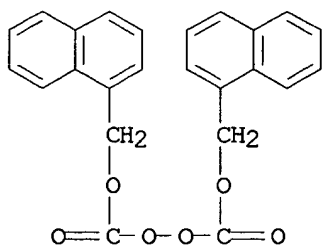
IT 138556-70-4 138556-73-7

RL: USES (Uses)

(catalyst-inhibitors, regulation of vinyl chloride radical polymn. in  
relation to)

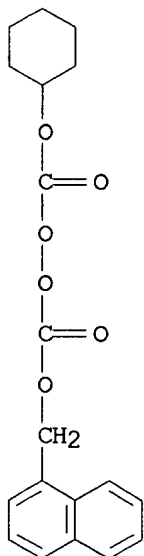
RN 138556-70-4 CAPLUS

CN Peroxydicarbonic acid, bis(1-naphthalenylmethyl) ester (9CI) (CA INDEX  
NAME)



RN 138556-73-7 CAPLUS

CN Peroxydicarbonic acid, cyclohexyl 1-naphthalenylmethyl ester (9CI) (CA  
INDEX NAME)



L13 ANSWER 7 OF 11 CAPLUS COPYRIGHT 2001 ACS

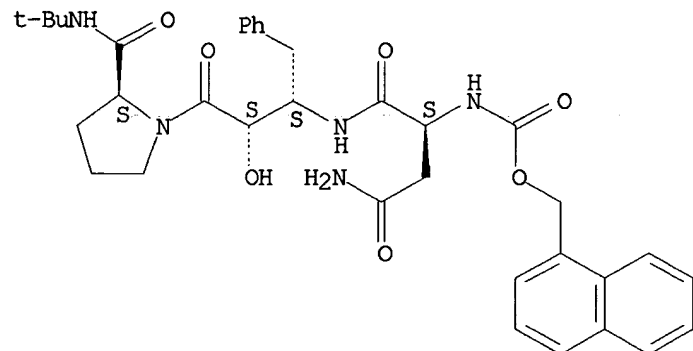
1993:204690 Document No. 118:204690 Kynostatin (KNI)-227 and -272, highly  
potent anti-HIV agents: conformationally constrained tripeptide  
inhibitors

of HIV protease containing allophenylnorstatine. Mimoto, Tsutomu; Imai,  
Junya; Kisanuki, Sumitsugu; Enomoto, Hiroshi; Hattori, Naoko; Akaji,  
Kenichi; Kiso, Yoshiaki (Dep. Med. Chem., Kyoto Pharm. Univ., Kyoto, 607,  
Japan). Chem. Pharm. Bull., 40(8), 2251-3 (English) 1992. CODEN:

CPBTAL.

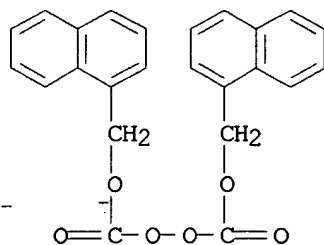
IT ISSN: 0009-2363.  
**143934-32-1**  
 RL: BIOL (Biological study)  
 (HIV protease inhibiting activity of, structure in relation to)  
 RN 143934-32-1 CAPLUS  
 CN L-Prolinamide, N2-[(1-naphthalenylmethoxy)carbonyl]-L-asparaginyl-  
 (.alpha.S,.beta.S)-.beta.-amino-.alpha.-hydroxybenzenebutanoyl-N-(1,1-  
 dimethylethyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

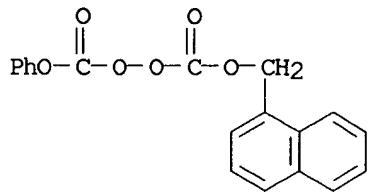


L13 ANSWER 8 OF 11 CAPLUS COPYRIGHT 2001 ACS  
 1992:613023 Document No. 117:213023 Initiator effect on late stages of  
 polymerization of vinyl chloride and methyl methacrylate. Grishin, A.  
 N.; Etlis, I. V.; Fomin, V. A.; Zegel'man, V. I.; Kulikova, G. L.; Radbil, T.  
 I.; Popov, V. A. (Nauchno-Issled. Inst. Khim. Tekhnol. Polim. im.  
 Kargina,  
 Dzerzhinsk, Russia). Vysokomol. Soedin., Ser. B, 34(6), 52-8 (Russian)  
 1992. CODEN: VYSBAI. ISSN: 0507-5483.

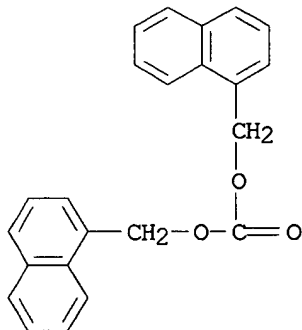
IT **138556-70-4 144255-52-7**  
 RL: CAT (Catalyst use); USES (Uses)  
 (catalysts, for radical polymn. of vinyl monomers, activity of,  
 structure in relation to)  
 RN 138556-70-4 CAPLUS  
 CN Peroxydicarbonic acid, bis(1-naphthalenylmethyl) ester (9CI) (CA INDEX  
 NAME)



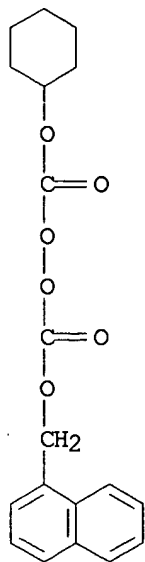
RN 144255-52-7 CAPLUS  
 CN Peroxydicarbonic acid, 1-naphthalenylmethyl phenyl ester (9CI) (CA INDEX  
 NAME)



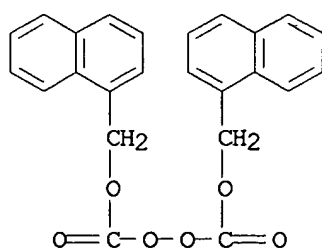
L13 ANSWER 9 OF 11 CAPLUS COPYRIGHT 2001 ACS  
 1992:213872 Document No. 116:213872 Synthesis and properties of peroxydicarbonates containing alkylaromatic fragments. Etlis, I. V.; Fomin, V. A.; Nozrina, F. D. (Nauchno-Issled. Inst. Khim.-Tekhnol. Polim., USSR). Zh. Org. Khim., 27(11), 2269-75 (Russian) 1991. CODEN: ZORKAE. ISSN: 0514-7492. OTHER SOURCES: CASREACT 116:213872.  
**IT 95225-95-9P 138556-73-7P**  
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)  
 (prepn. and thermolysis of, kinetics of)  
**RN 95225-95-9 CAPLUS**  
**CN 1-Naphthalenemethanol, carbonate (2:1) (9CI) (CA INDEX NAME)**



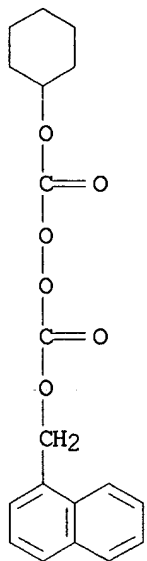
**RN 138556-73-7 CAPLUS**  
**CN Peroxydicarbonic acid, cyclohexyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)**



L13 ANSWER 10 OF 11 CAPLUS COPYRIGHT 2001 ACS  
 1992:42101 Document No. 116:42101 Polymerization of methyl methacrylate initiated with new alkylbenzyl (naphthylmethyl) peroxydicarbonates.  
 Etlis, I. V.; Fomin, V. A.; Radbil, T. I.; Malyshova, L. I.; Ovchinnikova,  
 Yu. I. (Nauchno-Issled. Inst. Khim. Tekhnol. Polimer. im. Kargina, USSR).  
 Vysokomol. Soedin., Ser. B, 33(9), 655-61 (Russian) 1991. CODEN: VYSBAI.  
 ISSN: 0507-5483.  
 IT 138556-70-4, Bis(1-naphthylmethyl) peroxydicarbonate  
 138556-73-7, 1-Naphthylmethyl cyclohexyl peroxydicarbonate  
 RL: CAT (Catalyst use); USES (Uses)  
 (catalysts, for radical polymn. of Me methacrylate, kinetics in relation to)  
 RN 138556-70-4 CAPLUS  
 CN Peroxydicarbonic acid, bis(1-naphthalenylmethyl) ester (9CI) (CA INDEX NAME)



RN 138556-73-7 CAPLUS  
 CN Peroxydicarbonic acid, cyclohexyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)



L13 ANSWER 11 OF 11 CAPLUS COPYRIGHT 2001 ACS  
1990:478951 Document No. 113:78951 Angiotensin-converting enzyme  
inhibitors:

synthesis and biological activity of N-substituted tripeptide inhibitors.  
Sawayama, Tadahiro; Tsukamoto, Masatoshi; Sasagawa, Takashi; Nishimura,  
Kazuya; Deguchi, Takashi; Takeyama, Kunihiko; Hosoki, Kanoo (Res. Lab.,  
Dainippon Pharm. Co., Ltd., Suita, 564, Japan). Chem. Pharm. Bull.,  
38(1), 110-15 (English) 1990. CODEN: CPBTAL. ISSN: 0009-2363. OTHER  
SOURCES: CASREACT 113:78951.

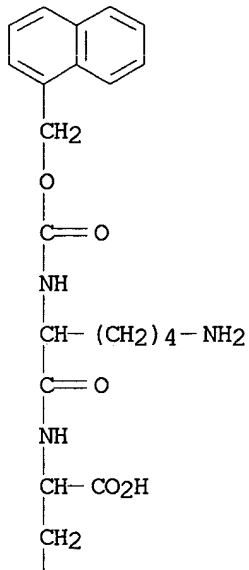
IT **116587-40-7P**

RL: SPN (Synthetic preparation); PREP (Preparation)  
(prepn. and inhibition by, of angiotensin-converting enzyme)

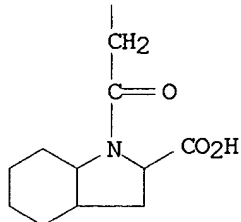
RN 116587-40-7 CAPLUS

CN D-Norvaline, 5-(2-carboxyoctahydro-1H-indol-1-yl)-N-[N2-[(1-naphthalenylmethoxy)carbonyl]-L-lysyl]-5-oxo-, [2S-(2.alpha.,3a.beta.,7a.beta.)]- (9CI) (CA INDEX NAME)

PAGE 1-A

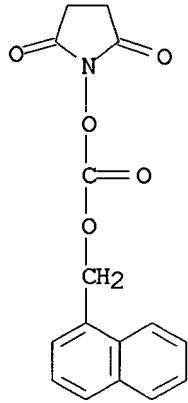


PAGE 2-A



IT 128595-04-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)  
(prepn. and sequential acylation of, with lysine tripeptide deriv.,  
and  
deblocking of, with trifluoroacetic acid)-  
RN 128595-04-0 CAPLUS  
CN 2,5-Pyrrolidinedione, 1-[[[1-naphthalenylmethoxy]carbonyl]oxy]- (9CI)  
(CA  
INDEX NAME)



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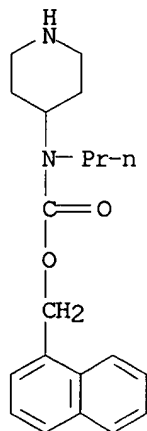
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L12 ANSWER 1 OF 32 CAPLUS COPYRIGHT 2001 ACS  
 2000:900455 Document No. 134:56574 Preparation of  
 aminopiperidinylmethylcyclopentanes as modulators of CCR-5 and/or CCR-3  
 chemokine receptors. Finke, Paul E.; Chapman, Kevin T.; Maccoss,  
 Malcolm;

Mills, Sander G.; Oates, Bryan (Merck & Co., Inc., USA). PCT Int. Appl.  
 WO 2000076512 A1 20001221, 223 pp. DESIGNATED STATES: W: AE, AG, AL,  
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AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ,  
 EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR,  
 KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ,  
 PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US,  
 UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF,  
 BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU,  
 MC, ML, MR, NE, NL, PT, SE, SN, TD, TG. (English). CODEN: PIXXD2.  
 APPLICATION: WO 2000-US15755 20000608. PRIORITY: US 1999-PV139067  
 19990611.

|    | PATENT NO.   | KIND   | DATE     | APPLICATION NO. | DATE     |
|----|--|--|----------|-----------------|----------|
| PI | WO 2000076512  | A1   | 20001221 | WO 2000-US15755 | 20000608 |
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|    | RW:  | GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG   |          |                 |          |
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|    | RL:  | RCT (Reactant)   |          |                 |          |
|    |  | (prepn. of aminopiperidinylmethylcyclopentanes as modulators of CCR-5 and/or CCR-3 chemokine receptors)  |          |                 |          |
| RN | 313527-12-7  | CAPLUS   |          |                 |          |
| CN | Carbamic acid, 4-piperidinylpropyl-, 1-naphthalenylmethyl ester, monohydrochloride (9CI) (CA INDEX NAME) |  |          |                 |          |



● HCl

L12 ANSWER 2 OF 32 CAPLUS COPYRIGHT 2001 ACS  
 2000:772600 Document No. 133:335461 Preparation and use of  
 2,4-diamino-3-hydroxy carboxylic acid derivatives as proteasome  
 inhibitors. France, Dennis; Furst, Peter; Zimmermann, Johann;  
 Garcia-Echeverria, Carlos; Scholz, Dieter; Furet, Pascal; Imbach,  
 Patricia  
 (Novartis A.-G., Switz.; Novartis-Erfindungen Verwaltungsgesellschaft  
 m.b.H.). PCT Int. Appl. WO 2000064863 A1 20001102, 38 pp. DESIGNATED  
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 RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI,  
 FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG.  
 (English). CODEN: PIXXD2. APPLICATION: WO 2000-EP3688 20000425.

PRIORITY: US 1999-300779 19990427; US 1999-388700 19990902.

PATENT NO. KIND DATE APPLICATION NO. DATE

PI WO 2000064863 A1 20001102 WO 2000-EP3688 20000425  
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SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA,  
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RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,  
DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,  
CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

IT 303186-89-2P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. and use of diaminohydroxy carboxylic acid derivs. as proteasome inhibitors)

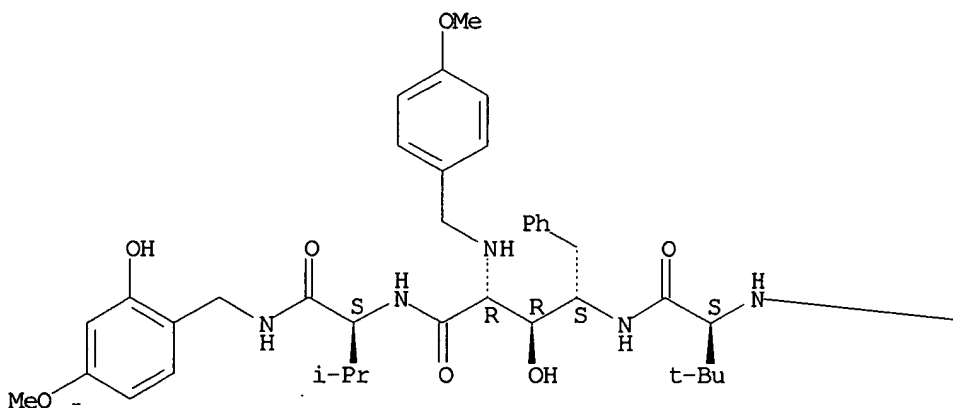
RN 303186-89-2 CAPLUS

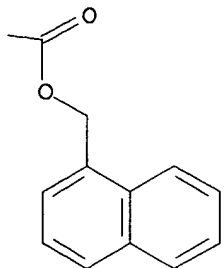
CN L-Valinamide,

N4-[3-methyl-N-[(1-naphthalenylmethoxy)carbonyl]-L-valyl]-4-amino-2,4,5-trideoxy-2-[[[(4-methoxyphenyl)methyl]amino]-5-phenyl-L-lyxonoyl-N-[(2-hydroxy-4-methoxyphenyl)methyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A





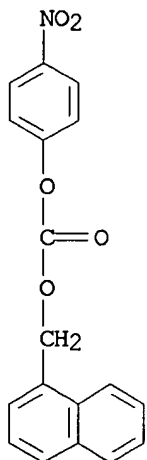
IT 172154-18-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)  
(prepn. and use of diaminohydroxy carboxylic acid derivs. as  
proteasome

inhibitors)

RN 172154-18-6 CAPLUS

CN Carbonic acid, 1-naphthalenylmethyl 4-nitrophenyl ester (9CI) (CA INDEX  
NAME)

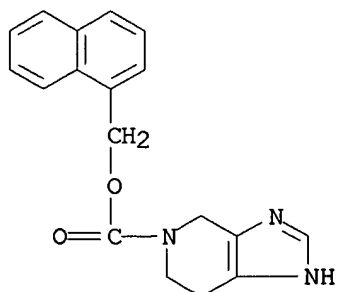


L12 ANSWER 3 OF 32 "CAPLUS" COPYRIGHT 2001 ACS

2000:756706 Document No. 133:321882 Preparation of substituted fused imidazoles for treatment and/or prevention of diseases and disorders related to the histamine H3 receptor. Dorwald, Florencio Zaragoza; Andersen, Knud Erik; Jorgensen, Tine Krogh; Peschke, Bernd; Wulff, Birgitte Schjellerup; Pettersson, Ingrid; Rudolf, Klaus; Stenkamp, Dirk; Hurnaus, Rudolf; Muller, Stephan Georg; Krist, Bernd (Novo Nordisk A/S,

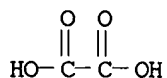
Den.; Boehringer Ingelheim International, G.m.b.H.). PCT Int. Appl. WO 2000063208 A1 20001026, 169 pp. DESIGNATED STATES: W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG. (English). CODEN: PIXXD2. APPLICATION: WO 2000-DK179 20000413. PRIORITY: DK 1999-508 19990416; DK 1999-1345 19990922; DK 2000-42 20000112.

|     | PATENT NO.   | KIND   | DATE     | APPLICATION NO.   | DATE     |
|-----|--|--------|----------|---|----------|
| PI  | WO 2000063208  | A1     | 20001026 | WO 2000-DK179   | 20000413 |
|     |  |        |          | W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM<br>RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG |          |
| IT  | 303020-31-7P   |        |          | RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)   |          |
|     |  |        |          | (prepn. of substituted fused imidazoles for treatment and/or prevention of diseases and disorders related to the histamine H3 receptor)   |          |
| RN  | 303020-31-7  | CAPLUS |          |   |          |
| CN  | 5H-Imidazo[4,5-c]pyridine-5-carboxylic acid, 1,4,6,7-tetrahydro-, 1-naphthalenylmethyl ester, ethanedioate (1:1) (9CI) (CA INDEX NAME) |        |          |   |          |
| CM  | 1  |        |          |   |          |
| CRN | 303020-30-6  |        |          |   |          |
| CMF | C18 H17 N3 O2  |        |          |   |          |



CM 2

CRN 144-62-7  
CMF C2 H2 O4

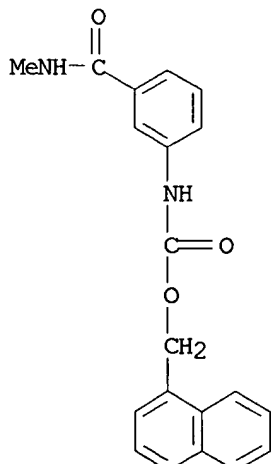


L12 ANSWER 4 OF 32 CAPLUS COPYRIGHT 2001 ACS  
 2000:441766 Document No. 133:43321 Amide derivatives. Ando, Ryoichi;  
 Chiba,

Noriko (Mitsubishi Chemical Corporation, Japan). PCT Int. Appl. WO  
 2000037434 A1 20000629, 46 pp. DESIGNATED STATES: W: AE, AL, AM, AT,  
 AU,

AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI,  
 GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,  
 LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO,  
 RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA,  
 ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH,  
 CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE,  
 NL, PT, SE, SN, TD, TG. (Japanese). CODEN: PIXXD2. APPLICATION: WO  
 1999-JP7138 19991220. PRIORITY: JP 1998-364499 19981222.

|    | PATENT NO.   | KIND            | DATE     | APPLICATION NO.   | DATE     |
|----|--|-----------------|----------|---|----------|
| PI | WO 2000037434  | A1              | 20000629 | WO 1999-JP7138  | 19991220 |
|    | W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,<br>CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,<br>IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA,<br>MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI,<br>SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ,<br>BY, KG, KZ, MD, RU, TJ, TM<br>RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,<br>DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,<br>CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG |                 |          |   |          |
| IT | <b>276252-43-8P</b>  |                 |          | RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic<br>preparation); THU (Therapeutic use); BIOL (Biological study); PREP<br>(Preparation); USES (Uses)<br>(prepn. of benzamides as antibacterial agents) |          |
| RN | 276252-43-8  | CAPLUS          |          |   |          |
| CN | Carbamic acid, [3-[(methylamino)carbonyl]phenyl]-, 1-naphthalenylmethyl<br>ester (9CI)   | (CA INDEX NAME) |          |   |          |

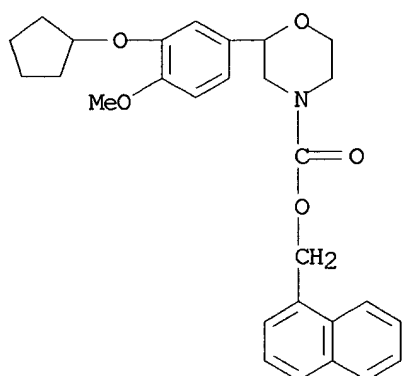


L12 ANSWER 5 OF 32 CAPLUS COPYRIGHT 2001 ACS

1999:748650 Document No. 132:12315 Preparation of 2-phenylmorpholine derivatives as phosphodiesterase inhibitors. Akiyama, Toshihiko; Ine, Shinji; Yamana, Kenjirou; Takahama, Akane (Nikken Chemicals Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 11322730 A2 19991124 Heisei, 49 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1999-59696 19990308.

PRIORITY: JP 1998-73059 19980309.

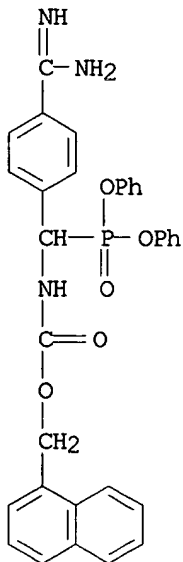
|    | PATENT NO.  | KIND            | DATE     | APPLICATION NO. | DATE     |
|----|---|-----------------|----------|-----------------|----------|
| PI | JP 11322730   | A2              | 19991124 | JP 1999-59696   | 19990308 |
| IT | <b>251315-16-9P</b>   |                 |          |                 |          |
|    | RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) |                 |          |                 |          |
|    | (prepn. of 2-phenylmorpholine derivs. as phosphodiesterase inhibitors)  |                 |          |                 |          |
| RN | 251315-16-9   | CAPLUS          |          |                 |          |
| CN | 4-Morpholinocarboxylic acid, 2-[3-(cyclopentyloxy)-4-methoxyphenyl]-, 1-naphthalenylmethyl ester (9CI)  | (CA INDEX NAME) |          |                 |          |



L12 ANSWER 6 OF 32 CAPLUS COPYRIGHT 2001 ACS

1999:582644 Document No. 131:214554 Preparation of basic .alpha.-aminoalkylphosphonate derivatives as serine protease inhibitors. Powers, James C.; Jackson, Delwin S.; Ni, Liming (Georgia Tech Research Corp., USA). U.S. US 5952307 A 19990914, 18 pp., Cont.-in-part of U.S. 5,686,419. (English). CODEN: USXXAM. APPLICATION: US 1997-907840-19970814. PRIORITY: US 1994-184286 19940121.

|    | PATENT NO.  | KIND            | DATE     | APPLICATION NO. | DATE     |
|----|---|-----------------|----------|-----------------|----------|
| PI | US 5952307  | A               | 19990914 | US 1997-907840  | 19970814 |
|    | US 5686419  | A               | 19971111 | US 1994-184286  | 19940121 |
| IT | <b>242816-96-2P</b>   |                 |          |                 |          |
|    | RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) |                 |          |                 |          |
|    | (prepn. of basic .alpha.-aminoalkylphosphonate derivs. as serine protease inhibitors)   |                 |          |                 |          |
| RN | 242816-96-2   | CAPLUS          |          |                 |          |
| CN | Carbamic acid, [[4-(aminoiminomethyl)phenyl](diphenoxypyrophosphinyl)methyl]-, 1-naphthalenylmethyl ester (9CI)   | (CA INDEX NAME) |          |                 |          |



L12 ANSWER 7 OF 32 CAPLUS COPYRIGHT 2001 ACS

1999:495290 Document No. 131:129991 [(Acylpyrrolo)methyl]imidazoles and analogs as farnesyl transferase inhibitors. Shin, You Seung; Koh, Jong Sung; Lee, Hyun Il; Lee, Jin Ho; Kim, Jong Hyun; Chung, Hyun Ho; Kim, Kwi Hwa; Kwak, Tae Hwan; Ro, Seong Gu; Ahn, In Ae; Choi, Tae Saeng; Oh, Young Hoon; Kim, Chung Mi; Lee, Sun Hwa; Kim, Hyun Sung (LG Chemical Ltd., S. Korea). PCT Int. Appl. WO 9938862 A1 19990805, 99 pp. DESIGNATED

STATES:

W: AU, BR, CA, CN, JP, MX, US; RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE. (English). CODEN: PIXXD2.

APPLICATION: WO 1999-KR51 19990201. PRIORITY: KR 1998-2776 19980202; KR 1998-2777 19980202; KR 1998-28340 19980714; KR 1998-32150 19980807.

PATENT NO. KIND DATE APPLICATION NO. DATE

| PI | WO 9938862  | A1 | 19990805 | WO 1999-KR51   | 19990201 |
|----|---|----|----------|----------------|----------|
|    | W: AU, BR, CA, CN, JP, MX, US<br>RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE |    |          |                |          |
|    | AU 9921886  | A1 | 19990816 | AU 1999-21886  | 19990201 |
|    | EP 1058683  | A1 | 20001213 | EP 1999-901979 | 19990201 |
|    | R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI                                   |    |          |                |          |

IT 234445-22-8P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of [(acylpvrrolo)methyl]imidazoles and analogs as farnesyl transferase inhibitors for treatment or prevention of cancer, restenosis, atherosclerosis, or infections from hepatitis delta and related diseases)

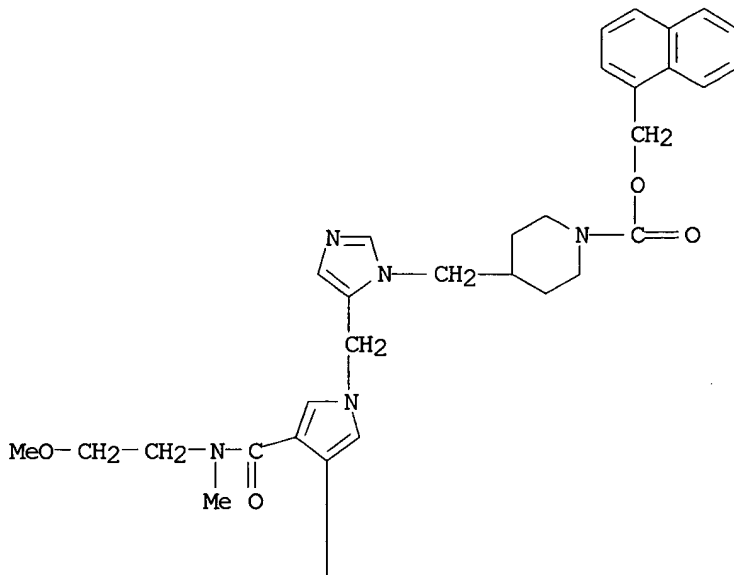
RN 234445-22-8 CAPLUS

CN 1-Piperidinecarboxylic acid,

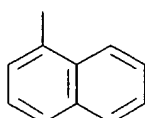
4-[[5-[[3-[(2-methoxyethyl)methylamino]carbo  
nyl]-4-(1-naphthalenyl)-1H-pyrrol-1-yl]methyl]-1H-imidazol-1-yl]methyl]-,

1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A



L12 ANSWER 8 OF 32 CAPLUS COPYRIGHT 2001 ACS

1999:96124 Document No. 130:168242 Preparation of 1-(4-sulfonamidobutyl)piperidines and related compounds as modulators of chemokine receptor activity.. Caldwell, Charles G.; Finke, Paul E.; MacCoss, Malcolm; Meurer, Laura C.; Mills, Sander G.; Oates, Bryan (Merck & Co., Inc., USA). PCT Int. Appl. WO 9904794 A1 19990204, 281 pp.

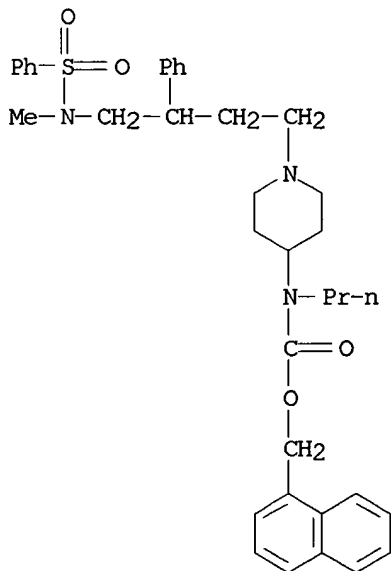
DESIGNATED STATES: W: AL, AM, AU, AZ, BA, BB, BG, BR, BY, CA, CN, CU, CZ,

EE, GE, HR, HU, ID, IL, IS, JP, KG, KR, KZ, LC, LK, LR, LT, LV, MD, MG, MK, MN, MX, NO, NZ, PL, RO, RU, SG, SI, SK, SL, TJ, TM, TR, TT, UA, US, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG. (English). CODEN: PIXXD2. APPLICATION: WO 1998-US14990 19980721. PRIORITY: US 1997-53754 19970725; GB 1998-958 19980116.

PATENT NO. KIND DATE APPLICATION NO. DATE

|    |  |    |          |                 |          |
|----|--|----|----------|-----------------|----------|
| PI | WO 9904794   | A1 | 19990204 | WO 1998-US14990 | 19980721 |
|    | W: AL, AM, AU, AZ, BA, BB, BG, BR, BY, CA, CN, CU, CZ, EE, GE, HR, HU, ID, IL, IS, JP, KG, KR, KZ, LC, LK, LR, LT, LV, MD, MG, MK, MN, MX, NO, NZ, PL, RO, RU, SG, SI, SK, SL, TJ, TM, TR, TT, UA, |    |          |                 |          |

US, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM  
 RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES,  
 FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI,  
 CM, GA, GN, GW, ML, MR, NE, SN, TD, TG  
 AU 9885760 A1 19990216 AU 1998-85760 19980721  
 EP 1003514 A1 20000531 EP 1998-936920 19980721  
 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, NL, SE, MC, PT, IE,  
 FI US 6136827 A 20001024 US 1998-120010 19980721  
 IT **220394-34-3P**  
 RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)  
 (prepn. of 1-(4-sulfonamidobutyl)piperidines and related compds. as modulators of chemokine receptor activity)  
 RN 220394-34-3 CAPLUS  
 CN Carbamic acid, [1-[4-[methyl(phenylsulfonyl)amino]-3-phenylbutyl]-4-piperidinyl]propyl-, 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)



L12 ANSWER 9 OF 32 CAPLUS COPYRIGHT 2001 ACS

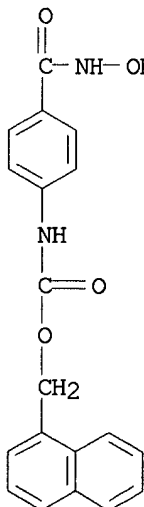
1997:752925 Document No. 128:34588 Preparation of benzohydroxamic acids as antiinflammatory and immunosuppressive agents.. Bertolini, Giorgio; Biffi, Mauro; Leoni, Flavio; Mizrahi, Jacques; Pavich, Gianfranco; Mascagni, Paolo (Italfarmaco S.P.A., Italy; Bertolini, Giorgio; Biffi, Mauro; Leoni, Flavio; Mizrahi, Jacques; Pavich, Gianfranco; Mascagni, Paolo). PCT Int. Appl. WO 9743251 A1 19971120, 44 pp. DESIGNATED

STATES:

W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, DE, DK, ES, FI, FR, GA, GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG.  
 (English). CODEN: PIXXD2. APPLICATION: WO 1997-EP2407 19970512.

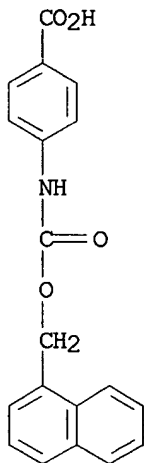
PRIORITY: IT 1996-MI968 19960514.

|    | PATENT NO.  | KIND | DATE     | APPLICATION NO. | DATE     |
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| PI | WO 9743251  | A1   | 19971120 | WO 1997-EP2407  | 19970512 |
|    | W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM |      |          |                 |          |
|    | RW: GH, KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG  |      |          |                 |          |
|    | CA 2254066  | AA   | 19971120 | CA 1997-2254066 | 19970512 |
|    | AU 9728964  | A1   | 19971205 | AU 1997-28964   | 19970512 |
|    | AU 713300   | B2   | 19991125 |                 |          |
|    | EP 901465   | A1   | 19990317 | EP 1997-923053  | 19970512 |
|    | EP 901465   | B1   | 20000927 |                 |          |
|    | R: DE, DK, ES, FR, GB, GR, NL, SE, PT, IE   |      |          |                 |          |
|    | CN 1221403  | A    | 19990630 | CN 1997-195410  | 19970512 |
|    | BR 9709234  | A    | 19990810 | BR 1997-9234    | 19970512 |
|    | JP 2000510472   | T2   | 20000815 | JP 1997-540505  | 19970512 |
|    | ES 2151267  | T3   | 20001216 | ES 1997-923053  | 19970512 |
|    | US 6034096  | A    | 20000307 | US 1998-180606  | 19981112 |
| IT | <b>199657-21-1P</b>   |      |          |                 |          |
|    | RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)   |      |          |                 |          |
|    | (prepn. of benzohydroxamic acids as antiinflammatory and immunosuppressive agents)  |      |          |                 |          |
| RN | 199657-21-1 CAPLUS  |      |          |                 |          |
| CN | Carbamic acid, [4-[(hydroxyamino)carbonyl]phenyl]-, 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)  |      |          |                 |          |



|    |  |
|----|--|
| IT | <b>199657-38-0P</b>  |
|    | RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)                |
|    | (prepn. of benzohydroxamic acids as antiinflammatory and immunosuppressive agents) |
| RN | 199657-38-0 CAPLUS   |
| CN | Benzoic acid, 4-[[[(1-naphthalenylmethoxy)carbonyl]amino]- (9CI) (CA INDEX         |

NAME)



L12 ANSWER 10 OF 32 CAPLUS COPYRIGHT 2001 ACS

1997:667724 Document No. 127:307384 Preparation of 3-

[(dioxoimidazolidinoacetyl)amino]-L-alanines and analogs as vitronectin receptor antagonists. Wehner, Volkmar; Knolle, Jochen; Stilz, Hans Ulrich; Carniato, Denis; Gourvest, Jean-Francois; Gadek, Tom; McDowell, Robert (Hoechst A.-G., Germany). Eur. Pat. Appl. EP 796855 A1 19970924, 115 pp. DESIGNATED STATES: R: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE. (German). CODEN: EPXXDW. APPLICATION: EP 1997-103712 19970306. PRIORITY: DE 1996-19610919 19960320; DE 1996-19626701 19960703; DE 1996-19635522 19960902.

| PATENT NO.  | KIND | DATE     | APPLICATION NO.  | DATE     |
|---|------|----------|------------------|----------|
| PI EP 796855  | A1   | 19970924 | EP 1997-103712   | 19970306 |
| SE R: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LI, LU, NL, PT, |      |          |                  |          |
| DE 19626701   | A1   | 19980108 | DE 1996-19626701 | 19960703 |
| DE 19635522   | A1   | 19980305 | DE 1996-19635522 | 19960902 |
| CA 2199923  | AA   | 19970920 | CA 1997-2199923  | 19970313 |
| AU 9716380  | A1   | 19970925 | AU 1997-16380    | 19970318 |
| AU 715729   | B2   | 20000210 |                  |          |
| NO 9701268  | A    | 19970922 | NO 1997-1268     | 19970319 |
| JP 09255664   | A2   | 19970930 | JP 1997-84711    | 19970319 |
| BR 9701335  | A    | 19980818 | BR 1997-1335     | 19970319 |
| ZA 9702381  | A    | 19981221 | ZA 1997-2381     | 19970319 |
| US 6218415  | B1   | 20010417 | US 1997-821253   | 19970320 |

IT 197357-96-3P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

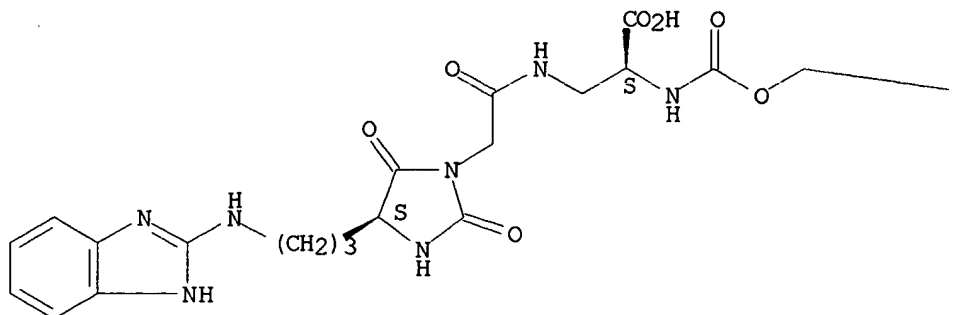
(prepn. of 3-[(dioxoimidazolidinoacetyl)amino]-L-alanines and analogs as vitronectin receptor antagonists)

RN 197357-96-3 CAPLUS

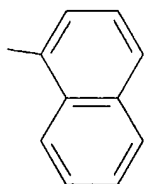
CN L-Alanine, 3-[[[(4S)-4-[3-(1H-benzimidazol-2-ylamino)propyl]-2,5-dioxo-1-imidazolidinyl]acetyl]amino]-N-[(1-naphthalenylmethoxy)carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A



PAGE 1-B



L12 ANSWER 11 OF 32 CAPLUS COPYRIGHT 2001 ACS

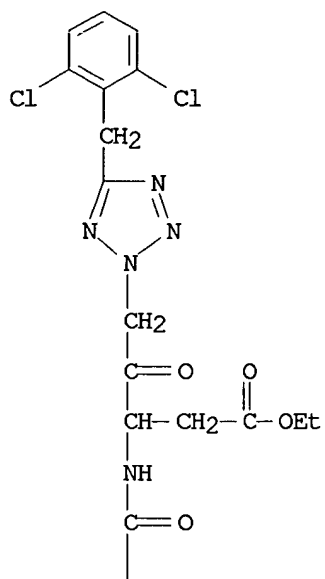
1997:491643 Document No. 127:109196 Preparation of tetrazole moiety-containing peptides as interleukin 1.beta. converting enzyme inhibitors. Ohmoto, Kazuyuki; Tanaka, Makoto; Miyazaki, Tohru; Ohno, Hiroyuki (Ono Pharmaceutical Co., Ltd., Japan; Ohmoto, Kazuyuki; Tanaka, Makoto; Miyazaki, Tohru; Ohno, Hiroyuki). PCT Int. Appl. WO 9724339 A1 19970710, 743 pp. DESIGNATED STATES: W: JP, KR, US; RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE. (Japanese). CODEN: PIXXD2. APPLICATION: WO 1996-JP3801 19961226. PRIORITY: JP 1995-351241 19951227.

|    | PATENT NO.  | KIND | DATE     | APPLICATION NO. | DATE     |
|----|---|------|----------|-----------------|----------|
| PI | WO 9724339  | A1   | 19970710 | WO 1996-JP3801  | 19961226 |
|    | W: JP, KR, US   |      |          |                 |          |
|    | RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, |      |          |                 |          |
| SE | EP 889039   | A1   | 19990107 | EP 1996-942651  | 19961226 |
|    | R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,  |      |          |                 |          |
|    | IE, FI  |      |          |                 |          |
|    | US 6136834  | A    | 20001024 | US 1998-101004  | 19980629 |

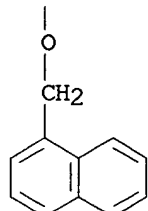
IT 192458-79-0P 192458-99-4P  
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)  
(prepn. of tetrazole moiety-contg. peptides as interleukin 1.beta. converting enzyme inhibitors)

RN 192458-79-0 CAPLUS  
CN 2H-Tetrazole-2-pentanoic acid,  
5-[(2,6-dichlorophenyl)methyl].beta.-[[[(1-naphthalenylmethoxy)carbonyl]amino]-.gamma.-oxo-, ethyl ester (9CI) (CA INDEX NAME)

PAGE 1-A



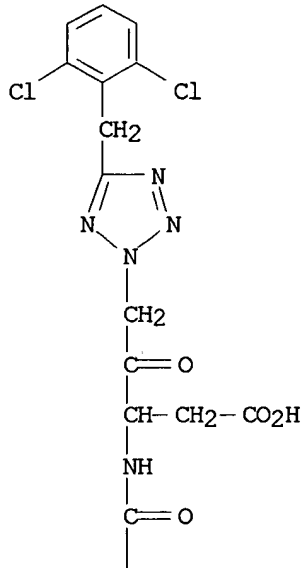
PAGE 2-A



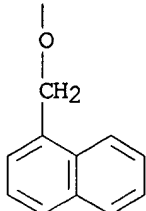
RN 192458-99-4 CAPLUS

CN 2H-Tetrazole-2-pentanoic acid,  
5-[(2,6-dichlorophenyl)methyl]-.beta.-[[[(1-  
naphthalenylmethoxy)carbonyl]amino]-.gamma.-oxo- (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A



L12 ANSWER 12 OF 32 CAPLUS COPYRIGHT 2001 ACS

1997:240459 Document No. 126:226021 Imidazole derivative catalysts for hardenable epoxy resin compositions. Nishikubo, Tatatomi (Taiyo Ink Mfg Co Ltd, Japan; Nishikubo Tatatomi). Jpn. Kokai Tokkyo Koho JP 09040750

A2

19970210 Heisei, 11 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP  
1996-149720 19960522. PRIORITY: JP 1995-146889 19950523.

PATENT NO. KIND DATE APPLICATION NO. DATE

| PI | JP 09040750 | A2 | 19970210 | JP 1996-149720 | 19960522 |
|----|-------------|----|----------|----------------|----------|
|    | US 5623023  | A  | 19970422 | US 1996-650981 | 19960521 |

IT 172359-58-9P

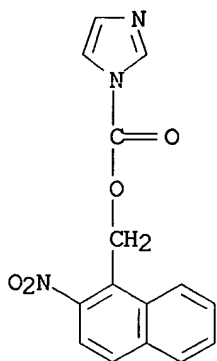
RL: CAT (Catalyst use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses)

(imidazole deriv. catalysts for hardenable epoxy resin compns.)

RN 172359-58-9 CAPLUS

CN 1H-Imidazole-1-carboxylic acid, (2-nitro-1-naphthalenyl)methyl ester (9CI)

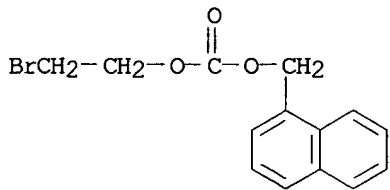
(CA INDEX NAME)



L12 ANSWER 13 OF 32 CAPLUS COPYRIGHT 2001 ACS  
 1996:134049 Document No. 124:175810 Preparation of heterocyclic compounds as

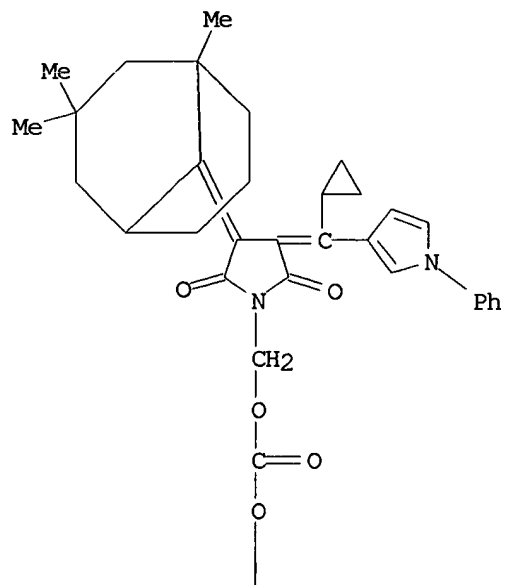
photochromic substances. Tanizawa, Tsuneyoshi; Kobayakawa, Takashi (Tokuyama Kk, Japan). Jpn. Kokai Tokkyo Koho JP 07285931 A2 19951031 Heisei, 35 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1994-80685 19940419.

|    | PATENT NO.  | KIND | DATE     | APPLICATION NO. | DATE     |
|----|---|------|----------|-----------------|----------|
| PI | JP 07285931   | A2   | 19951031 | JP 1994-80685   | 19940419 |
| IT | <b>123498-61-3</b>  |      |          |                 |          |
| RL | RCT (Reactant)<br>(prepn. of heterocyclic compds. as photochromic substances) |      |          |                 |          |
| RN | 123498-61-3 CAPLUS  |      |          |                 |          |
| CN | Carbonic acid, 2-bromoethyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)  |      |          |                 |          |

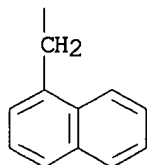


IT **173972-38-8P**  
 RL: SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)  
 (prepn. of heterocyclic compds. as photochromic substances)  
 RN 173972-38-8 CAPLUS  
 CN Carbonic acid, [3-[cyclopropyl(1-phenyl-1H-pyrrol-3-yl)methylene]-2,5-dioxo-4-(1,3,3-trimethylbicyclo[3.3.1]non-9-ylidene)-1-pyrrolidinyl]methyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A



L12 ANSWER 14 OF 32 CAPLUS COPYRIGHT 2001 ACS

1995:974107 Document No. 124:89021 Thiocarbonate curing agents and curable polymer compositions containing them. Nishikubo, Tatatomi; Kameyama, Atsushi; Narita, Kichihei; Hagio, Shigeru; Uehara, Shinichi (San Nopco

Kk,

Japan). Jpn. Kokai Tokkyo Koho JP 07252212 A2 19951003 Heisei, 6 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1994-67937 19940312.

PATENT NO. KIND DATE APPLICATION NO. DATE

PI JP 07252212 A2 19951003 JP 1994-67937 19940312

IT 172359-58-9

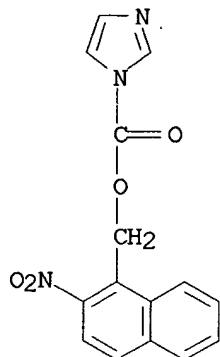
RL: RCT (Reactant)

(in prepn. of thiocarbonate curing agents for photocurable polymer compns.)

RN 172359-58-9 CAPLUS

CN 1H-Imidazole-1-carboxylic acid, (2-nitro-1-naphthalenyl)methyl ester (9CI)

(CA INDEX NAME)

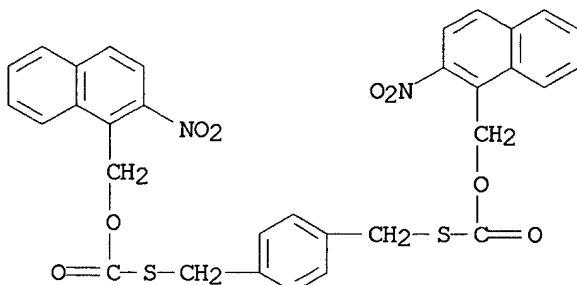


IT 172359-55-6P, p-Xylenebis(2-nitro-.alpha.-naphthalenemethyl-.alpha.-S-thiocarbonate)

RL: MOA (Modifier or additive use); PNU (Preparation, unclassified); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (thiocarbonate curing agents and photocurable polymer compns. with good storage stability)

RN 172359-55-6 CAPLUS

CN Carbonothioic acid, S,S'-[1,4-phenylenebis(methylene)]O,O'-bis[(2-nitro-1-naphthalenyl)methyl] ester (9CI) (CA INDEX NAME)



L12 ANSWER 15 OF 32 CAPLUS COPYRIGHT 2001 ACS

1995:875007 Document No. 124:55952 Preparation of N-substituted (3R,4R)-3-ethyl[(1-methyl-1H-imidazol-5-yl)methyl]-2-pyrrolidone antiglaucoma agents. Albaugh, Pamela; White, Gregory J.; Garst, Michael E. (Allergan, Inc., USA). U.S. US 5453434 A 19950926, 6 pp. Cont.-in-part of U.S. Ser. No. 126,285. (English). CODEN: USXXAM.

APPLICATION: US 1994-265163 19940624. PRIORITY: US 1989-434929 19891113; US 1993-126285 19930920.

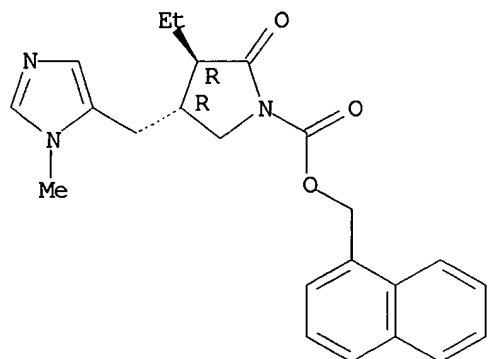
| PATENT NO.    | KIND  | DATE     | APPLICATION NO. | DATE     |
|---------------|-------|----------|-----------------|----------|
| -----         | ----- | -----    | -----           | -----    |
| PI US 5453434 | A     | 19950926 | US 1994-265163  | 19940624 |
| US 5264449    | A     | 19931123 | US 1989-434929  | 19891113 |

IT 172154-25-5P

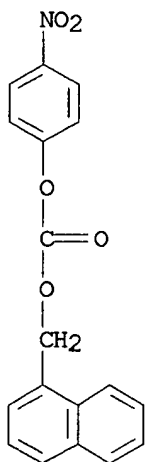
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of N-substituted (3R,4R)-3-ethyl[(1-methyl-1H-imidazol-5-yl)methyl]-2-pyrrolidone antiglaucoma agents)

RN 172154-25-5 CAPLUS  
CN 1-Pyrrolidinecarboxylic acid, 3-ethyl-4-[(1-methyl-1H-imidazol-5-yl)methyl]-2-oxo-, 1-naphthalenylmethyl ester, (3R-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



IT 172154-18-6  
RL: RCT (Reactant)  
(prepn. of N-substituted (3R,4R)-3-ethyl[(1-methyl-1H-imidazol-5-yl)methyl]-2-pyrrolidone antiglaucoma agents)  
RN 172154-18-6 CAPLUS  
CN Carbonic acid, 1-naphthalenylmethyl 4-nitrophenyl ester (9CI) (CA INDEX NAME)



L12 ANSWER 16 OF 32 CAPLUS COPYRIGHT 2001 ACS  
1995:520421 Document No. 122:265356 Preparation of fulgide and fulgimide photochromic compounds. Imura, Tomohito; Tanizawa, Tsuneyoshi; Kobayakawa, Takashi (Tokuyama Soda Kk, Japan). Jpn. Kokai Tokkyo Koho JP 06345772 A2 19941220 Heisei, 8 pp. (Japanese). CODEN: JKXXAF.  
APPLICATION: JP 1993-167315 19930615.

| PATENT NO.     | KIND | DATE     | APPLICATION NO. | DATE     |
|----------------|------|----------|-----------------|----------|
| PI JP 06345772 | A2   | 19941220 | JP 1993-167315  | 19930615 |

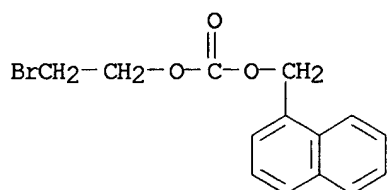
IT 123498-61-3

RL: RCT (Reactant)

(prepn. of fulgide and fulgimide photochromic compds.)

RN 123498-61-3 CAPLUS

CN Carbonic acid, 2-bromoethyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)



IT 162689-53-4P

RL: SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)

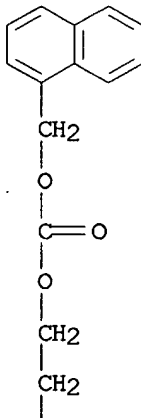
(prepn. of fulgide and fulgimide photochromic compds.)

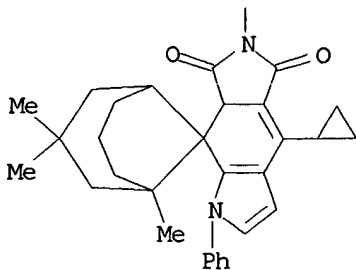
RN 162689-53-4 CAPLUS

CN Carbonic acid,

2-(4-cyclopropyl-1,5,7,7a-tetrahydro-1',3',3'-trimethyl-5,7-dioxo-1-phenylspiro[benzo[1,2-b:4,5-c']dipyrrole-8(6H),9'-bicyclo[3.3.1]nonan]-6-yl)ethyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)

PAGE 1-A

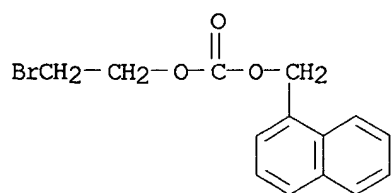




L12 ANSWER 17 OF 32 CAPLUS COPYRIGHT 2001 ACS

1995:520378 Document No. 122:265237 Preparation of spirofulgide and -fulgimide analogs as photochromic compounds. Imura, Satoshi; Tanizawa, Tsuneyoshi; Kobayakawa, Takashi (Tokuyama Corp., Japan). Eur. Pat. Appl. EP 629626 A2 19941221, 69 pp. DESIGNATED STATES: R: DE, ES, FR, IT. (English). CODEN: EPXXDW. APPLICATION: EP 1994-304140 19940608. PRIORITY: JP 1993-141023 19930611.

| PATENT NO.  | KIND | DATE     | APPLICATION NO. | DATE     |
|---|------|----------|-----------------|----------|
| PI EP 629626  | A2   | 19941221 | EP 1994-304140  | 19940608 |
| EP 629626   | A3   | 19950301 |                 |          |
| EP 629626   | B1   | 19991027 |                 |          |
| R: DE, ES, FR, IT   |      |          |                 |          |
| JP 07002824   | A2   | 19950106 | JP 1993-141023  | 19930611 |
| JP 3138117  | B2   | 20010226 |                 |          |
| AU 9464634  | A1   | 19941215 | AU 1994-64634   | 19940608 |
| AU 679513   | B2   | 19970703 |                 |          |
| ES 2140506  | T3   | 20000301 | ES 1994-304140  | 19940608 |
| US 5708063  | A    | 19980113 | US 1996-601832  | 19960215 |
| IT 123498-61-3, Carbonic acid, 2-bromoethyl 1-naphthalenylmethyl ester          |      |          |                 |          |
| RL: RCT (Reactant)  |      |          |                 |          |
| (prepn. of spirofulgide and -fulgimide analogs as photochromic compds.)         |      |          |                 |          |
| RN 123498-61-3 CAPLUS   |      |          |                 |          |
| CN Carbonic acid, 2-bromoethyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME) |      |          |                 |          |



IT 162689-53-4P

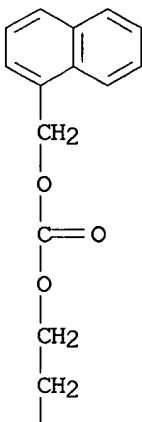
RL: SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)  
(prepn. of spirofulgide and -fulgimide analogs as photochromic compds.)

RN 162689-53-4 CAPLUS

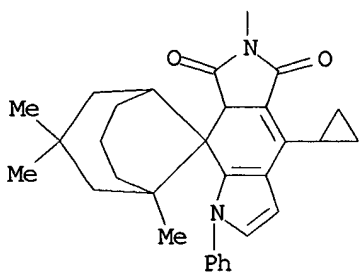
CN Carbonic acid,  
2-(4-cyclopropyl-1,5,7,7a-tetrahydro-1',3',3'-trimethyl-5,7-

dioxo-1-phenylspiro[benzo[1,2-b:4,5-c']dipyrrole-8(6H),9'-bicyclo[3.3.1]nonan]-6-yl)ethyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A



L12 ANSWER 18 OF 32 CAPLUS COPYRIGHT 2001 ACS

1995:330664 Document No. 122:105635 preparation of heterocyclyl-containing ketones as drugs. Ando, Ryoichi; Ando, Naoko; Masuda, Hirokazu; Sakaki, Toshiro; Morinaka, Yasuhiro; Takahashi, Chizuko; Tamao, Yoshikuni; Tobe, Akihiro (Mitsubishi Chem Ind, Japan). Jpn. Kokai Tokkyo Koho JP 06192199 A2 19940712 Heisei, 252 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1992-359273 19921225.

| PATENT NO.      | KIND  | DATE     | APPLICATION NO. | DATE     |
|-----------------|-------|----------|-----------------|----------|
| -----           | ----- | -----    | -----           | -----    |
| PI JP 06192199  | A2    | 19940712 | JP 1992-359273  | 19921225 |
| IT 160652-75-5P |       |          |                 |          |

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

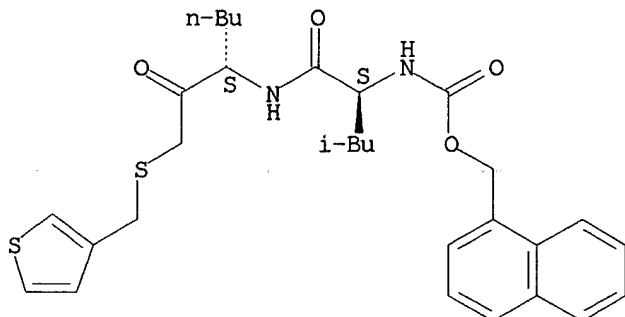
(prepn. of heterocyclyl-contg. ketones as drugs)

RN 160652-75-5 CAPLUS

CN Carbamic acid,

[3-methyl-1-[[[1-[(3-thienylmethyl)thio]acetyl]pentyl]amin  
o]carbonylbutyl]-, 1-naphthalenylmethyl ester, [S-(R\*,R\*)]- (9CI) (CA  
INDEX NAME)

Absolute stereochemistry.



L12 ANSWER 19 OF 32 CAPLUS COPYRIGHT 2001 ACS

1995:299775 Document No. 122:82080 Preparation of analogs of cholecystokinin

(30-33) containing an .alpha.-substituted aminoacid as drugs. Horwell, David Christopher; Howson, William; Hugues, John; Richardson, Reginald Stewart (Warner-Lambert Co., USA). PCT Int. Appl. WO 9409031 A1 19940428,

71 pp. DESIGNATED STATES: W: AU, CA, CZ, FI, HU, JP, KR, NO, NZ, RU, SK;

RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE.  
(English). CODEN: PIXXD2. APPLICATION: WO 1993-US9809 19931014.

PRIORITY: US 1992-963169 19921019; US 1993-131693 19931008.

PATENT NO. KIND DATE APPLICATION NO. DATE

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PI WO 9409031 A1 19940428 WO 1993-US9809 19931014  
W: AU, CA, CZ, FI, HU, JP, KR, NO, NZ, RU, SK  
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE  
AU 9453596 A1 19940509 AU 1994-53596 19931014

IT 146034-77-7P 146034-78-8P 146034-79-9P

146034-82-4P 146034-83-5P 160280-21-7P

160280-22-8P 160280-23-9P 160280-24-0P

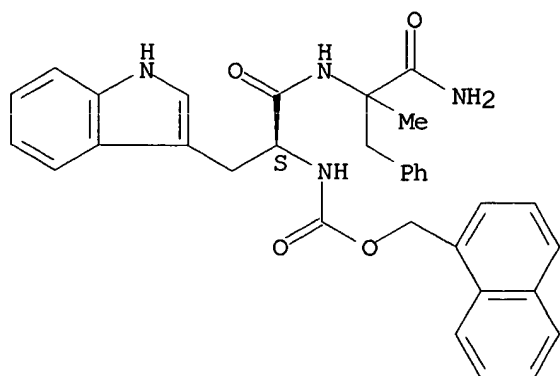
160280-25-1P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(prepn. of, as cholecystokinin analog)

RN 146034-77-7 CAPLUS

CN Phenylalaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl- (9CI) (CA INDEX NAME)

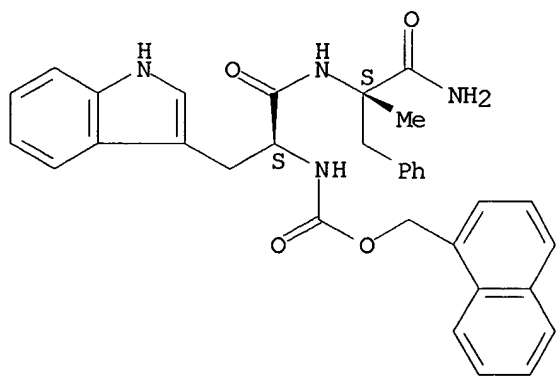
Absolute stereochemistry.



RN 146034-78-8 CAPLUS

CN L-Phenylalaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl- (9CI) (CA INDEX NAME)

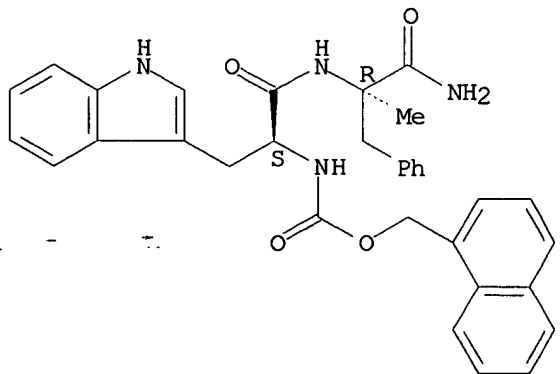
Absolute stereochemistry.



RN 146034-79-9 CAPLUS

CN D-Phenylalaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl- (9CI) (CA INDEX NAME)

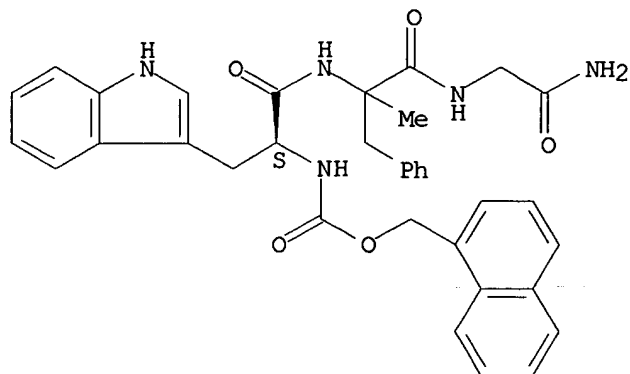
Absolute stereochemistry.



RN 146034-82-4 CAPLUS

CN Glycinamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methylphenylalanyl- (9CI) (CA INDEX NAME)

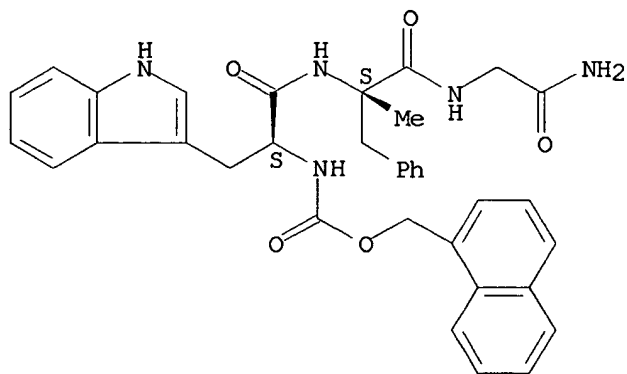
Absolute stereochemistry.



RN 146034-83-5 CAPLUS

CN Glycinamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl-L-phenylalanyl- (9CI) (CA INDEX NAME)

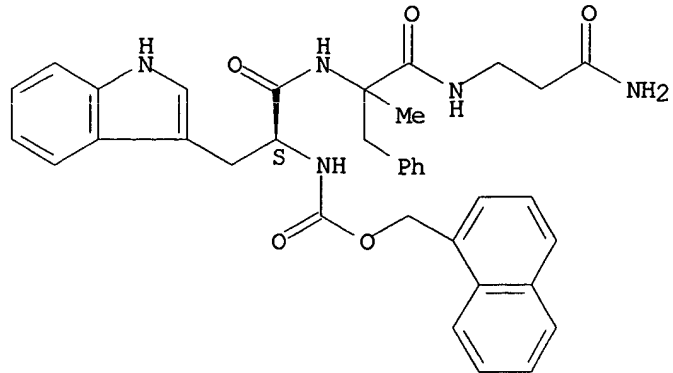
Absolute stereochemistry.



RN 160280-21-7 CAPLUS

CN .beta.-Alaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methylphenylalanyl- (9CI) (CA INDEX NAME)

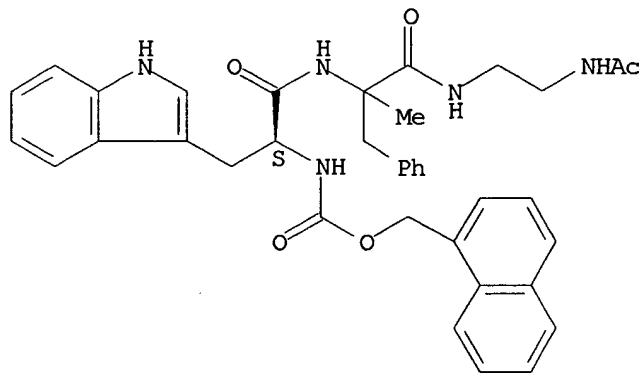
Absolute stereochemistry.



RN 160280-22-8 CAPLUS

CN Phenylalaninamide, N-[ (1-naphthalenylmethoxy) carbonyl]-L-tryptophyl-N-[2-(acetylamino)ethyl]-.alpha.-methyl- (9CI) (CA INDEX NAME)

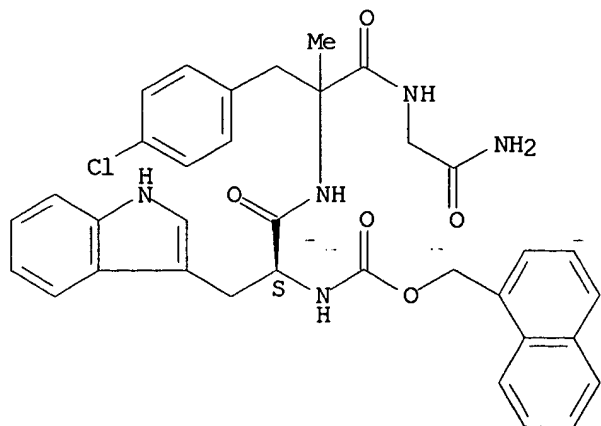
Absolute stereochemistry.



RN 160280-23-9 CAPLUS

CN Glycinamide, N-[ (1-naphthalenylmethoxy) carbonyl]-L-tryptophyl-4-chloro-.alpha.-methylphenylalanyl- (9CI) (CA INDEX NAME)

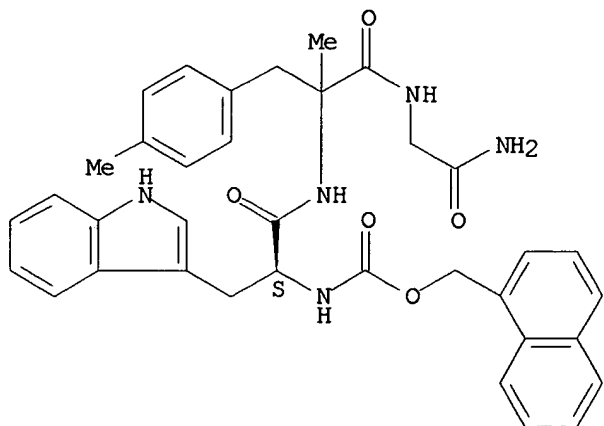
Absolute stereochemistry.



RN 160280-24-0 CAPLUS

CN Glycinamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.,4-dimethylphenylalanyl- (9CI) (CA INDEX NAME)

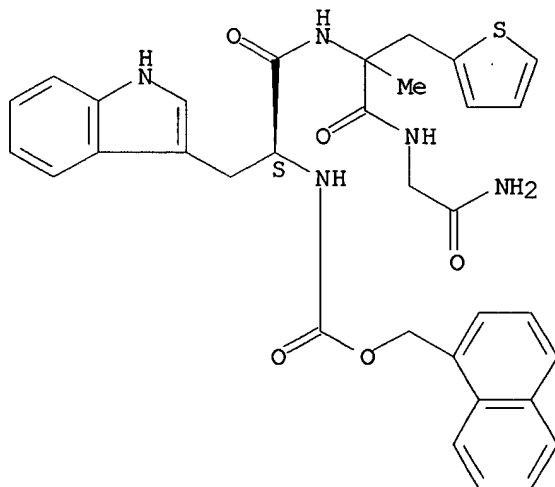
Absolute stereochemistry.



RN 160280-25-1 CAPLUS

CN Glycinamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-2-methyl-3-(2-thienyl)alanyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



IT 160280-27-3P 160280-28-4P 160280-30-8P

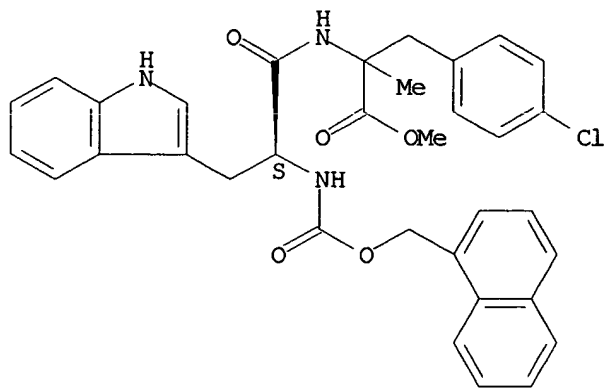
160280-31-9P 160280-34-2P 160280-35-3P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(prepn. of, as intermediate for cholecystokinin analog)

RN 160280-27-3 CAPLUS

CN Phenylalanine, 4-chloro-.alpha.-methyl-N-[N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl]-, methyl ester (9CI) (CA INDEX NAME)

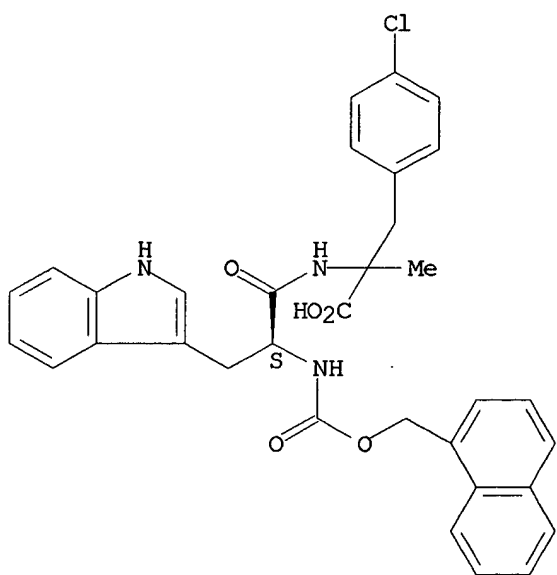
Absolute stereochemistry.



RN 160280-28-4 CAPLUS

CN Phenylalanine, 4-chloro-.alpha.-methyl-N-[N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl]- (9CI) (CA INDEX NAME)

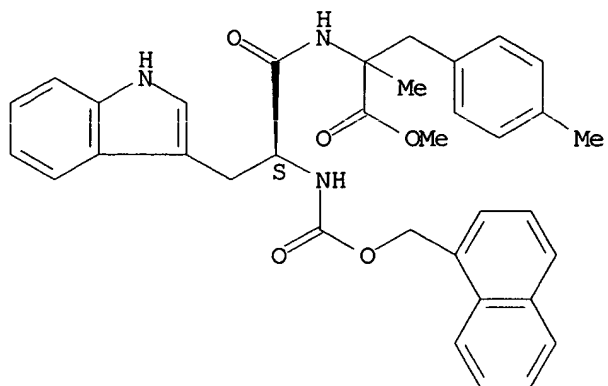
Absolute stereochemistry.



RN 160280-30-8 CAPLUS

CN Phenylalanine,  
.alpha.,4-dimethyl-N-[N-[(1-naphthalenylmethoxy)carbonyl]-L-  
tryptophyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

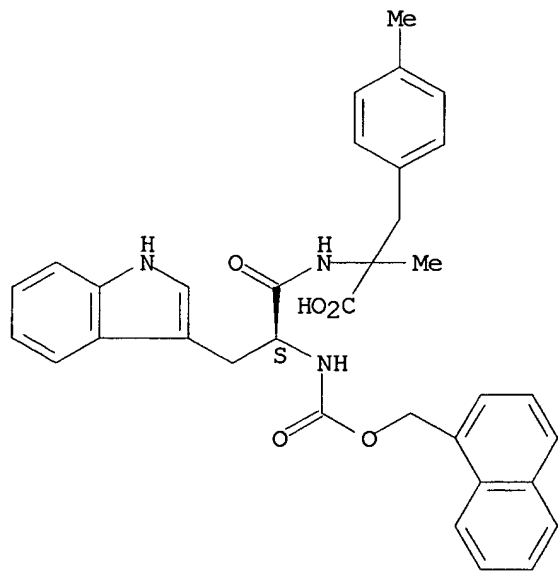


RN 160280-31-9 CAPLUS

CN Phenylalanine,

.alpha.,4-dimethyl-N-[N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl]- (9CI) (CA INDEX NAME)

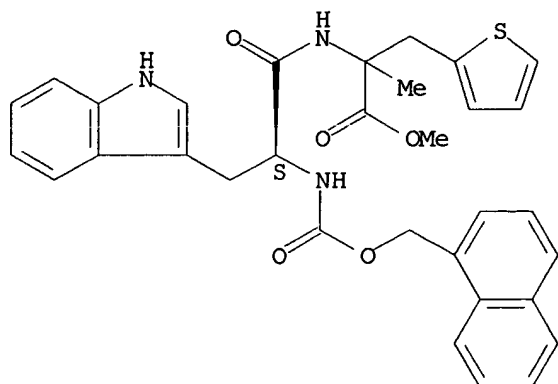
Absolute stereochemistry.



RN 160280-34-2 CAPLUS

CN Alanine, 2-methyl-N-[N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl]-3-(2-thienyl)-, methyl ester (9CI) (CA INDEX NAME)

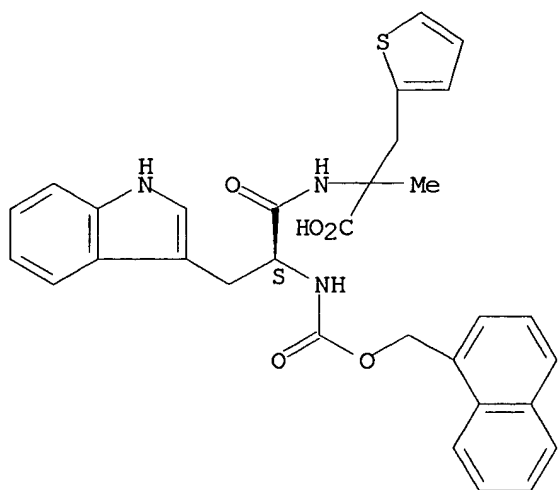
Absolute stereochemistry.



RN 160280-35-3 CAPLUS

CN Alanine, 2-methyl-N-[N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl]-3-(2-thienyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



IT 39545-08-9

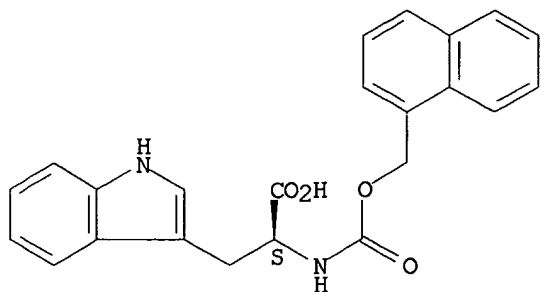
RL: RCT (Reactant)

(reaction of, in prepn. of cholecystokinin analog)

RN 39545-08-9 CAPLUS

CN L-Tryptophan, N-[(1-naphthalenylmethoxy)carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L12 ANSWER 20 OF 32 CAPLUS COPYRIGHT 2001 ACS  
 1993:409161 Document No. 119:9161 HIV protease inhibitors. Mimoto,  
 Tsutomu;  
 Hattori, Naoko; Nagano, Yuuichi; Shintani, Makoto; Kiso, Yoshiaki (Nippon  
 Mining Co., Ltd., Japan). Eur. Pat. Appl. EP 490667 A2 19920617, 86 pp.  
 DESIGNATED STATES: R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU,  
 NL,

SE. (English). CODEN: EPXXDW. APPLICATION: EP 1991-311549 19911211.  
 PRIORITY: JP 1990-409673 19901211; JP 1991-25755 19910125; JP 1991-89976  
 19910328; JP 1991-169174 19910614; JP 1991-304043 19911023.

|    | PATENT NO.  | KIND | DATE     | APPLICATION NO. | DATE     |
|----|---|------|----------|-----------------|----------|
| PI | EP 490667   | A2   | 19920617 | EP 1991-311549  | 19911211 |
|    | EP 490667   | A3   | 19930505 |                 |          |
|    | EP 490667   | B1   | 19990609 |                 |          |
|    | R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE |      |          |                 |          |
|    | CA 2056911  | AA   | 19920612 | CA 1991-2056911 | 19911204 |
|    | CA 2056911  | C    | 19980922 |                 |          |
|    | JP 05170722   | A2   | 19930709 | JP 1991-348705  | 19911205 |
|    | JP 2700511  | B2   | 19980121 |                 |          |
|    | AU 9188900  | A1   | 19920618 | AU 1991-88900   | 19911206 |
|    | AU 653972   | B2   | 19941020 |                 |          |
|    | ZA 9109721  | A    | 19921230 | ZA 1991-9721    | 19911210 |
|    | FI 9105819  | A    | 19920612 | FI 1991-5819    | 19911211 |
|    | AT 181080   | E    | 19990615 | AT 1991-311549  | 19911211 |
|    | ES 2134764  | T3   | 19991016 | ES 1991-311549  | 19911211 |
|    | NO 9200023  | A    | 19920727 | NO 1992-23      | 19920102 |

IT 143934-32-1P 143934-40-1P 143934-54-7P

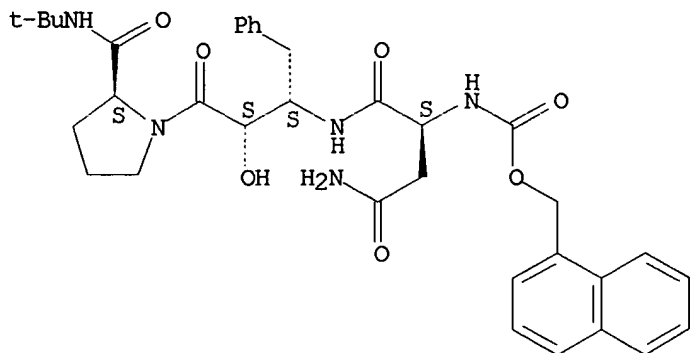
143934-89-8P

RL: SPN (Synthetic preparation); PREP (Preparation)  
 (prepn. and HIV protease-inhibiting activity of)

RN 143934-32-1 CAPLUS

CN L-Prolinamide, N2-[ (1-naphthalenylmethoxy)carbonyl]-L-asparaginyl-  
 (.alpha.S,.beta.S)-.beta.-amino-.alpha.-hydroxybenzenebutanoyl-N-(1,1-  
 dimethylethyl)- (9CI) (CA INDEX NAME)

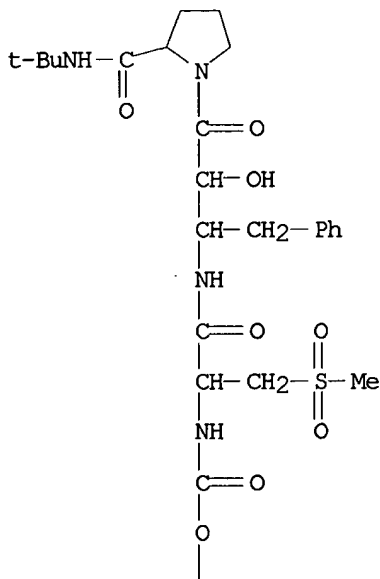
Absolute stereochemistry.



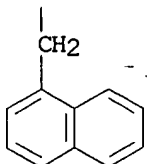
RN 143934-40-1 CAPLUS

CN L-Prolinamide, 3-(methylsulfonyl)-N-[(1-naphthalenylmethoxy)carbonyl]-L-alanyl-(2S,3S)-2-hydroxy-4-phenyl-3-aminobutanoyl-N-(1,1-dimethylethyl)-(9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A

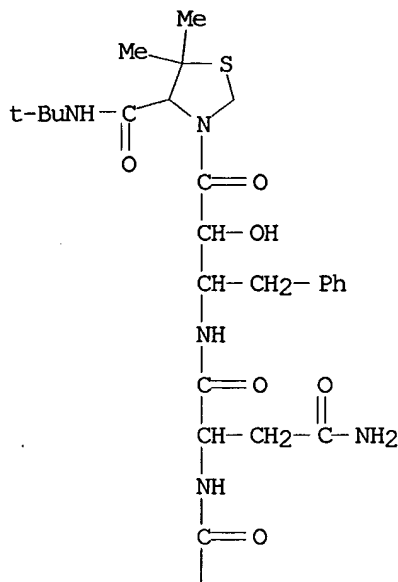


RN 143934-54-7 CAPLUS

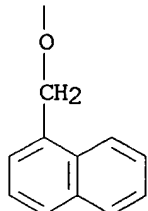
CN Carbamic acid, [3-amino-1-[[3-[4-[(1,1-dimethylethyl)amino]carbonyl]-5-

dimethyl-3-thiazolidinyl]-2-hydroxy-3-oxo-1-(phenylmethyl)propyl]amino]carbonyl]-3-oxopropyl]-, 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A



RN 143934-89-8 CAPLUS

CN L-Prolinamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-.alpha.-aspartyl-(2S,3S)-2-hydroxy-4-phenyl-3-aminobutanoyl-N-(1,1-dimethylethyl)-, hydrazide, monoacetate (salt) (9CI) (CA INDEX NAME)

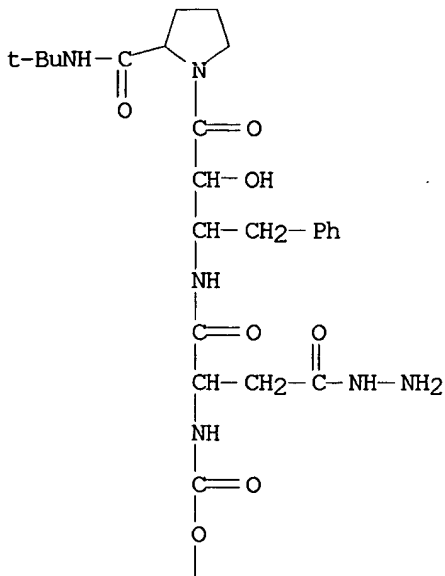
CM 1

CRN 143934-88-7

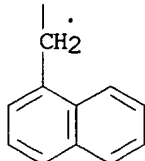
CMF C35 H44 N6 O7

CDES \*

PAGE 1-A

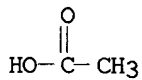


PAGE 2-A



CM 2

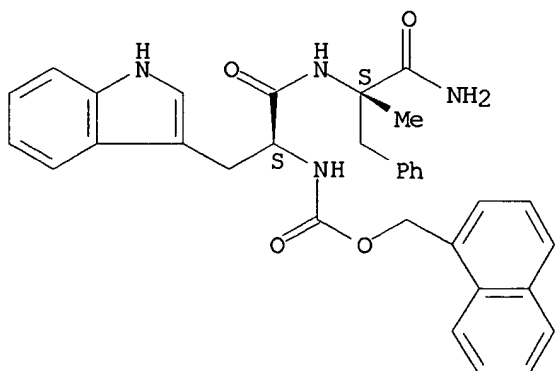
CRN 64-19-7  
CMF C2 H4 O2



L12 ANSWER 21 OF 32 CAPLUS COPYRIGHT 2001 ACS  
1993:94333 Document No. 118:94333 .alpha.-Substituted polypeptides having therapeutic activity. Horwell, David Christopher; Hugues, John; Richardson, Reginald Stewart; Howson, William (Warner-Lambert Co., USA). PCT Int. Appl. WO 9219254 A1 19921112, 45 pp. DESIGNATED STATES: W: AU, CA, JP; RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, MC, NL, SE. (English). CODEN: PIXXD2. APPLICATION: WO 1992-US3119 19920415. PRIORITY: US 1991-690755 19910424; US 1992-852086 19920320.

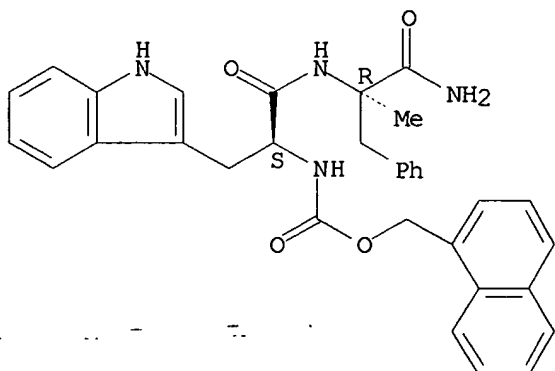
| PATENT NO.   | KIND | DATE     | APPLICATION NO. | DATE     |
|--|------|----------|-----------------|----------|
| PI WO 9219254  | A1   | 19921112 | WO 1992-US3119  | 19920415 |
| W: AU, CA, JP<br>RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, MC, NL, SE                                    |      |          |                 |          |
| AU 9219072   | A1   | 19921221 | AU 1992-19072   | 19920415 |
| JP 06507402  | T2   | 19940825 | JP 1992-511401  | 19920415 |
| EP 668770  | A1   | 19950830 | EP 1992-911434  | 19920415 |
| R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, MC, NL, SE  |      |          |                 |          |
| ZA 9202956   | A    | 19931025 | ZA 1992-2956    | 19920423 |
| IT 146034-78-8 146034-79-9 146034-83-5   |      |          |                 |          |
| RL: BIOL (Biological study)<br>(for analgesic or other therapeutic)  |      |          |                 |          |
| RN 146034-78-8 CAPLUS  |      |          |                 |          |
| CN L-Phenylalaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl- (9CI) (CA INDEX NAME) |      |          |                 |          |

Absolute stereochemistry.



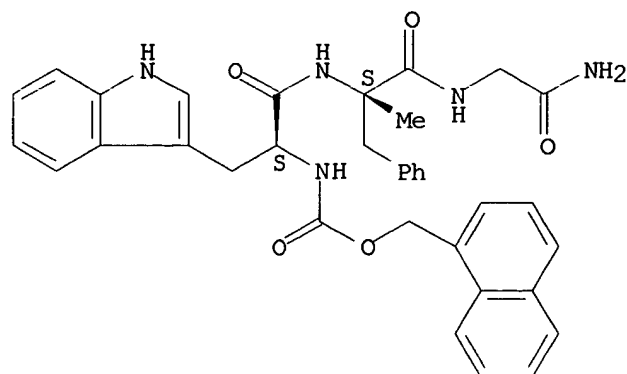
RN 146034-79-9 CAPLUS  
CN D-Phenylalaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



RN 146034-83-5 CAPLUS  
CN Glycinamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl-L-phenylalanyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



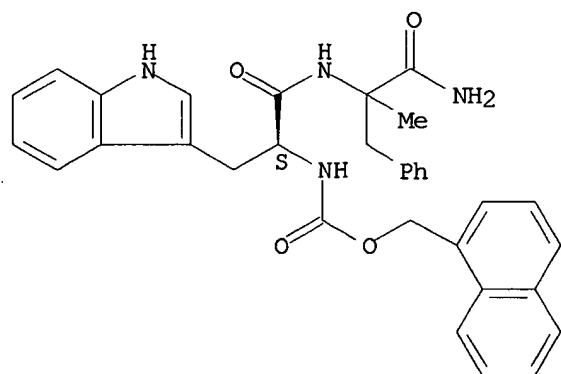
IT 146034-77-7P 146034-82-4P

RL: SPN (Synthetic preparation); PREP (Preparation)  
(prepn. of, for analgesic or other therapeutic)

RN 146034-77-7 CAPLUS

CN Phenylalaninamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methyl- (9CI) (CA INDEX NAME)

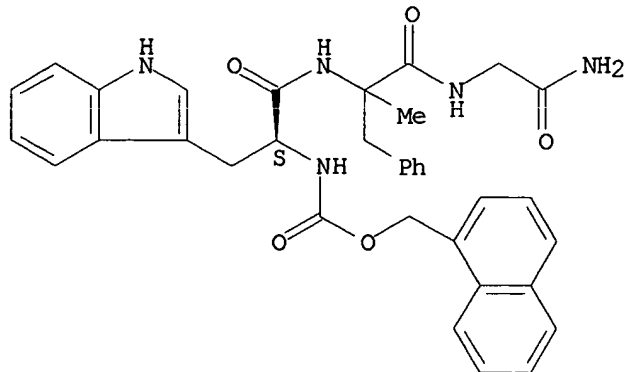
Absolute stereochemistry.



RN 146034-82-4 CAPLUS

CN Glycinamide, N-[(1-naphthalenylmethoxy)carbonyl]-L-tryptophyl-.alpha.-methylphenylalanyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



IT 39545-08-9

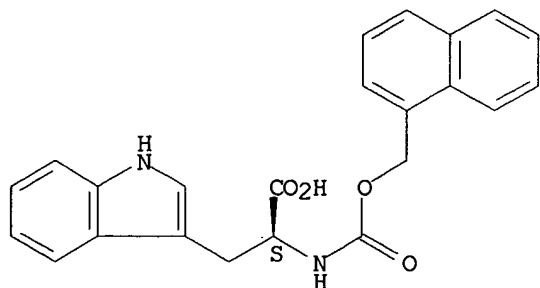
RL: RCT (Reactant)

(reaction of, for peptide prepn. for analgesic or other therapeutic)

RN 39545-08-9 CAPLUS

CN L-Tryptophan, N-[1-naphthalenylmethoxy]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L12 ANSWER 22 OF 32 CAPLUS COPYRIGHT 2001 ACS

1992:459004 Document No. 117:59004 Photochromic composition. Momota, Junji;

Kawaguchi, Ikuzo; Tanaka, Takashi; Kida, Yasuji (Tokuyama Soda Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 03121188 A2 19910523 Heisei, 33 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1989-257800 19891004.

PATENT NO. KIND DATE APPLICATION NO. DATE

| PI | JP 03121188 | A2 | 19910523 | JP 1989-257800 | 19891004 |
|----|-------------|----|----------|----------------|----------|
|    | JP 07033508 | B4 | 19950412 |                |          |

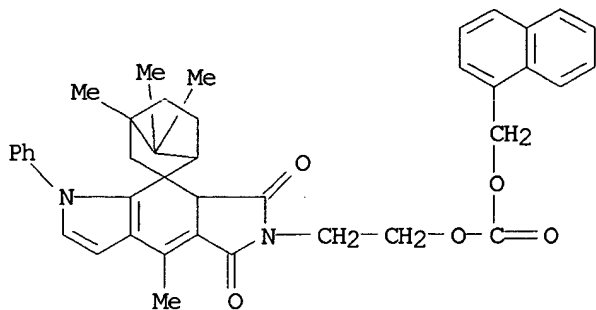
IT 123498-25-9

RL: USES (Uses)

(photochromic compn. contg.)

RN 123498-25-9 CAPLUS

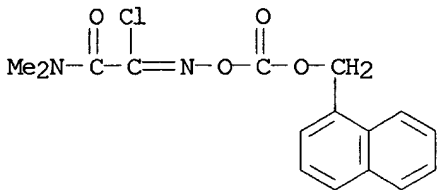
CN Carbonic acid, 1-naphthalenylmethyl 2-(1,5,7,7a-tetrahydro-4,4',7',7'-tetramethyl-5,7-dioxo-1-phenylspiro[benzo[1,2-b:4,5-c']dipyrrole-8(6H),2'bicyclo[2.2.1]heptan]-6-yl)ethyl ester (9CI) (CA INDEX NAME)



L12 ANSWER 23 OF 32 CAPLUS COPYRIGHT 2001 ACS

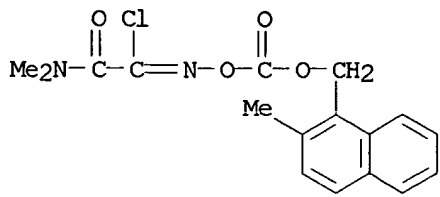
1992:407677 Document No. 117:7677 Oxime carbonates as fungicides. Adams, John Benjamin, Jr. (du Pont de Nemours, E. I., and Co., USA). PCT Int. Appl. WO 9204318 A1 19920319, 44 pp. DESIGNATED STATES: W: AU, BR, HU, JP, KR, SU, US; RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE. (English). CODEN: PIXXD2. APPLICATION: WO 1991-US5588 19910814. PRIORITY: US 1990-573073 19900829.

|    | PATENT NO.   | KIND                                | DATE   | APPLICATION NO. | DATE     |
|----|--|-------------------------------------|--|-----------------|----------|
| PI | WO 9204318   | A1                                  | 19920319   | WO 1991-US5588  | 19910814 |
|    |  | W:                                  | AU, BR, HU, JP, KR, SU, US                         |                 |          |
|    |  | RW:                                 | AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE |                 |          |
|    | AU 9184994   | A1                                  | 19920330   | AU 1991-84994   | 19910814 |
|    | CN 1059712   | A                                   | 19920325   | CN 1991-108591  | 19910829 |
| IT | 141700-15-4P 141700-19-8P 141700-22-3P   |                                     |  |                 |          |
|    | RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) |                                     |  |                 |          |
|    |  | (prepn. and fungicidal activity of) |  |                 |          |
| RN | 141700-15-4  | CAPLUS                              |  |                 |          |
| CN | Ethanimidoyl chloride, 2-(dimethylamino)-N-[(1-naphthalenylmethoxy)carbonyl]oxy]-2-oxo-  | (9CI)                               | (CA INDEX NAME)                                    |                 |          |



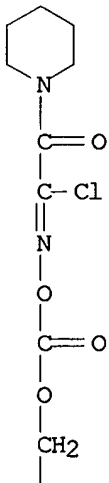
RN 141700-19-8 CAPLUS

CN Ethanimidoyl chloride, 2-(dimethylamino)-N-[[[(2-methyl-1-naphthalenyl)methoxy]carbonyl]oxy]-2-oxo- (9CI) (CA INDEX NAME)

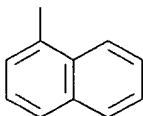


RN 141700-22-3 CAPLUS  
 CN 1-Piperidineethanimidoyl chloride,  
 N-[[ (1-naphthalenylmethoxy)carbonyl]oxy  
 ]-.alpha.-oxo- (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A

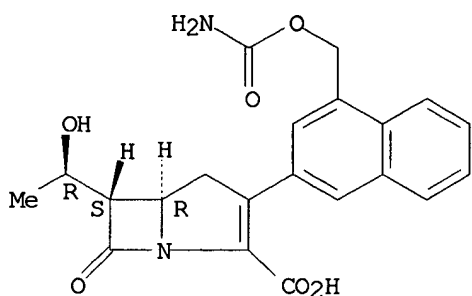


L12 ANSWER 24 OF 32 CAPLUS COPYRIGHT 2001 ACS  
 1992:255394 Document No. 116:255394 Preparation of 2-naphthyl-carbapenems.  
 Dininno, Frank P.; Greenlee, Mark L. (Merck and Co., Inc., USA). Eur.  
 Pat. Appl. EP 466253 A1 19920115, 59 pp. DESIGNATED STATES: R: CH, DE,  
 FR, GB, IT, LI, NL. (English). CODEN: EPXXDW. APPLICATION: EP  
 1991-201705 19910703. PRIORITY: US 1990-551707 19900711; US 1990-594510  
 19901009.

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|------|------|-----------------|------|
|------------|------|------|-----------------|------|

PI EP 466253 A1 19920115 EP 1991-201705 19910703  
 R: CH, DE, FR, GB, IT, LI, NL  
 US 5006519 A 19910409 US 1990-551707 19900711  
 US 5132422 A 19920721 US 1990-594510 19901009  
 IT 135869-05-5P  
 RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)  
 (prepn. of, as antibacterial)  
 RN 135869-05-5 CAPLUS  
 CN 1-Azabicyclo[3.2.0]hept-2-ene-2-carboxylic acid, 3-[4-  
 [[(aminocarbonyl)oxy]methyl]-2-naphthalenyl]-6-(1-hydroxyethyl)-7-oxo-,  
 monopotassium salt, [5R-[5.alpha.,6.alpha.(R\*)]]- (9CI) (CA INDEX NAME)

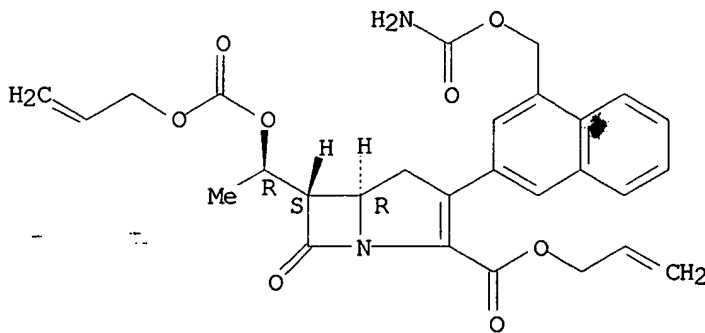
Absolute stereochemistry.



● K

IT 139768-15-3P 141433-48-9P  
 RL: SPN (Synthetic preparation); PREP (Preparation)  
 (prepn. of, as intermediate for naphthylcarbapenem antibacterial)  
 RN 139768-15-3 CAPLUS  
 CN 1-Azabicyclo[3.2.0]hept-2-ene-2-carboxylic acid, 3-[4-  
 [[(aminocarbonyl)oxy]methyl]-2-naphthalenyl]-7-oxo-6-[1-[(2-  
 propenoxy)carbonyl]oxy]ethyl-, 2-propenyl ester, [5R-  
 [5.alpha.,6.alpha.(R\*)]]- (9CI) (CA INDEX NAME)

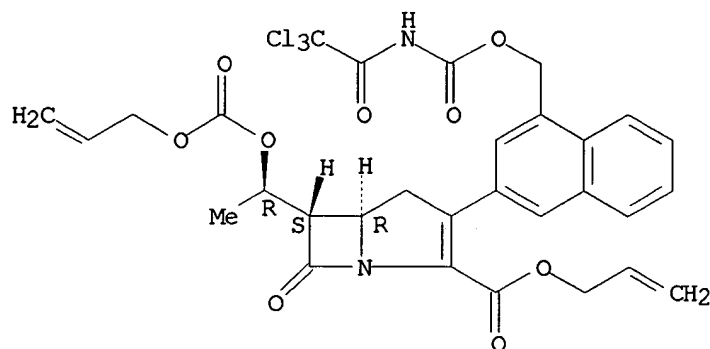
Absolute stereochemistry.



RN 141433-48-9 CAPLUS  
 CN 1-Azabicyclo[3.2.0]hept-2-ene-2-carboxylic acid, 7-oxo-6-[1-[(2-

propenyloxy)carbonyl]oxy]ethyl]-3-[4-[[[[trichloroacetyl]amino]carbonyl]oxy]methyl]-2-naphthalenyl]-, 2-propenyl ester, [5R-[5.alpha.,6.alpha.(R\*)]]- (9CI) (CA INDEX NAME)

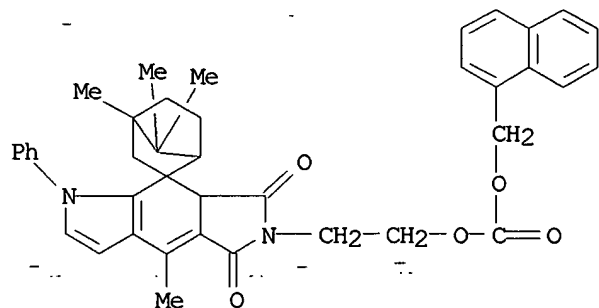
Absolute stereochemistry.



L12 ANSWER 25 OF 32 CAPLUS COPYRIGHT 2001 ACS

1992:140180 Document No. 116:140180 Composition of photochromic material.  
Momota, Junji; Kawaguchi, Ikuzo; Tanaka, Takashi; Kida, Yasuji (Tokuyama Soda Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 03124790 A2 19910528 Heisei, 23 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 1989-263001 19891011.

|    | PATENT NO.   | KIND | DATE     | APPLICATION NO. | DATE     |
|----|--|------|----------|-----------------|----------|
| PI | JP 03124790  | A2   | 19910528 | JP 1989-263001  | 19891011 |
|    | JP 07033509  | B4   | 19950412 |                 |          |
| IT | <b>123498-25-9P</b>  |      |          |                 |          |
|    | RL: SPN (Synthetic preparation); PREP (Preparation)<br>(prepn. and use of, photochromic material from)   |      |          |                 |          |
| RN | 123498-25-9 CAPLUS   |      |          |                 |          |
| CN | Carbonic acid, 1-naphthalenylmethyl 2-(1,5,7,7a-tetrahydro-4,4',7',7'-tetramethyl-5,7-dioxo-1-phenylspiro[benzo[1,2-b:4,5-c']dipyrrole-8(6H),2'-bicyclo[2.2.1]heptan]-6-yl)ethyl ester (9CI) (CA INDEX NAME) |      |          |                 |          |

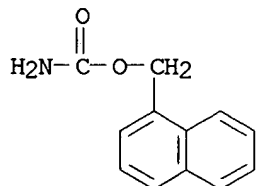


L12 ANSWER 26 OF 32 CAPLUS COPYRIGHT 2001 ACS

1992:53700 Document No. 116:53700 Supersorbent material as pesticide  
potentiator. Puritch, George S.; McHarg, Douglas; Bradbury, Roderick;

Mason, Wenda (Safer, Inc., USA). U.S. US 5037654 A 19910806, 7 pp.  
(English). CODEN: USXXAM. APPLICATION: US 1988-187589 19880428.

|    | PATENT NO.   | KIND | DATE     | APPLICATION NO. | DATE     |
|----|--|------|----------|-----------------|----------|
| PI | US 5037654   | A    | 19910806 | US 1988-187589  | 19880428 |
|    | CA 1330710   | A1   | 19940719 | CA 1989-588469  | 19890117 |
| IT | <b>74156-18-6</b>  |      |          |                 |          |
|    | RL: BIOL (Biological study)<br>(polyacrylamide as potentiator for) |      |          |                 |          |
| RN | 74156-18-6 CAPLUS  |      |          |                 |          |
| CN | 1-Naphthalenemethanol, carbamate (9CI) (CA INDEX NAME)             |      |          |                 |          |

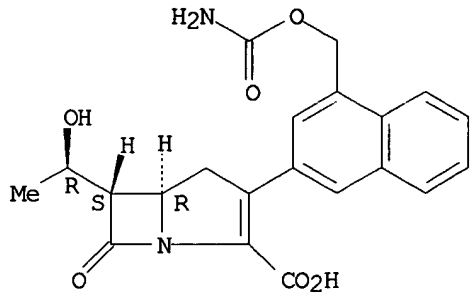


L12 ANSWER 27 OF 32 CAPLUS COPYRIGHT 2001 ACS

1991:558832 Document No. 115:158832 Preparation of 2-naphthylcarbapenem antibacterial agents. Dininno, Frank P.; Greenlee, Mark L. (Merck and Co., Inc., USA). U.S. US 5006519 A 19910409, 27 pp. (English). CODEN: USXXAM. APPLICATION: US 1990-551707 19900711.

|    | PATENT NO.  | KIND | DATE     | APPLICATION NO. | DATE     |
|----|---|------|----------|-----------------|----------|
| PI | US 5006519  | A    | 19910409 | US 1990-551707  | 19900711 |
|    | EP 466253   | A1   | 19920115 | EP 1991-201705  | 19910703 |
|    | R: CH, DE, FR, GB, IT, LI, NL   |      |          |                 |          |
|    | CA 2046505  | AA   | 19920112 | CA 1991-2046505 | 19910709 |
|    | JP 04230384   | A2   | 19920819 | JP 1991-171354  | 19910711 |
| IT | <b>135869-05-5P</b>   |      |          |                 |          |
|    | RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)<br>(prepn. of, as antibacterial)  |      |          |                 |          |
| RN | 135869-05-5 CAPLUS  |      |          |                 |          |
| CN | 1-Azabicyclo[3.2.0]hept-2-ene-2-carboxylic acid, 3-[4-<br>[(aminocarbonyl)oxy]methyl]-2-naphthalenyl]-6-(1-hydroxyethyl)-7-oxo-,<br>monopotassium salt, [5R-[5.alpha.,6.alpha.(R*)]]- (9CI) (CA INDEX NAME) |      |          |                 |          |

Absolute stereochemistry.

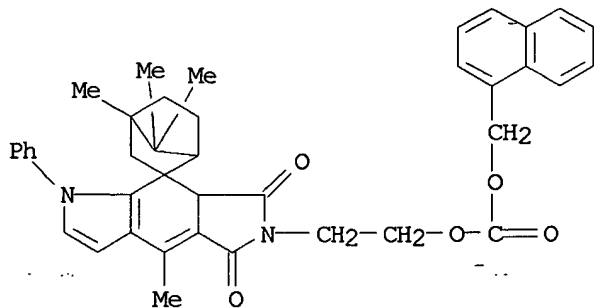


● K

L12 ANSWER 28 OF 32 CAPLUS COPYRIGHT 2001 ACS

1989:632775 Document No. 111:232775 Preparation of fused-ring fulgides and fulgimides as photochromic substances. Tanaka, Takashi; Imura, Satoshi; Kida, Yasuji (Tokuyama Soda Co., Ltd., Japan). Eur. Pat. Appl. EP 316179 A2 19890517, 98 pp. DESIGNATED STATES: R: DE, FR, IT. (English). CODEN: EPXXDW. APPLICATION: EP 1988-310608 19881110. PRIORITY: JP 1987-282131 19871110; JP 1987-283116 19871111; JP 1988-80250 19880402.

|    | PATENT NO.  | KIND | DATE     | APPLICATION NO. | DATE     |
|----|---|------|----------|-----------------|----------|
| PI | EP 316179   | A2   | 19890517 | EP 1988-310608  | 19881110 |
|    | EP 316179   | A3   | 19901212 |                 |          |
|    | EP 316179   | B1   | 19940119 |                 |          |
|    | R: DE, FR, IT   |      |          |                 |          |
|    | JP 01052778   | A2   | 19890228 | JP 1987-282131  | 19871110 |
| IT | <b>123498-25-9P</b>   |      |          |                 |          |
|    | RL: SPN (Synthetic preparation); PREP (Preparation)<br>(prepn. of, as photochromic substance)   |      |          |                 |          |
| RN | 123498-25-9 CAPLUS  |      |          |                 |          |
| CN | Carbonic acid, 1-naphthalenylmethyl 2-(1,5,7,7a-tetrahydro-4,4',7',7'-tetramethyl-5,7-dioxo-1-phenylspiro[benzo[1,2-b:4,5-c']dipyrrole-8(6H),2''-bicyclo[2.2.1]heptan]-6-yl)ethyl ester (9CI) (CA INDEX NAME) |      |          |                 |          |

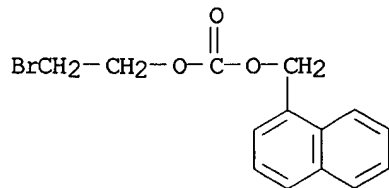


IT **123498-61-3**

RL: RCT (Reactant)  
(reaction of, in prepn. of photochromic substances)

RN 123498-61-3 CAPLUS

CN Carbonic acid, 2-bromoethyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME)

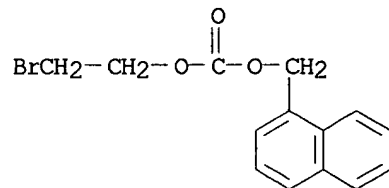


L12 ANSWER 29 OF 32 CAPLUS COPYRIGHT 2001 ACS

1989:605543 Document No. 111:205543 Fulgides as photochromic substances and a process for their preparation. Tanaka, Takashi; Imura, Tomohito; Kida, Yasuji (Tokuyama Soda Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 01052778 A2 19890228 Heisei, 27 pp. (Japanese). CODEN: JKXXAF.

APPLICATION: JP 1987-282131 19871110. PRIORITY: JP 1987-133370 19870530.

|    | PATENT NO.   | KIND   | DATE     | APPLICATION NO. | DATE     |
|----|--|--------|----------|-----------------|----------|
| PI | JP 01052778  | A2     | 19890228 | JP 1987-282131  | 19871110 |
|    | JP 02028154  | A2     | 19900130 | JP 1988-277495  | 19881104 |
|    | JP 07045502  | B4     | 19950517 |                 |          |
|    | US 4882438   | A      | 19891121 | US 1988-268497  | 19881108 |
|    | AU 8825005   | A1     | 19890511 | AU 1988-25005   | 19881110 |
|    | AU 615491  | B2     | 19911003 |                 |          |
|    | EP 316179  | A2     | 19890517 | EP 1988-310608  | 19881110 |
|    | EP 316179  | A3     | 19901212 |                 |          |
|    | EP 316179  | B1     | 19940119 |                 |          |
|    | R: DE, FR, IT  |        |          |                 |          |
|    | US 4960678   | A      | 19901002 | US 1989-403487  | 19890906 |
| IT | <b>123498-61-3</b> , 2-Bromoethyl 1-naphthylmethyl carbonate                 |        |          |                 |          |
|    | RL: RCT (Reactant)   |        |          |                 |          |
|    | (alkylation by, of furano-, thieno-, or pyrrolophthalimide deriv.)           |        |          |                 |          |
| RN | 123498-61-3  | CAPLUS |          |                 |          |
| CN | Carbonic acid, 2-bromoethyl 1-naphthalenylmethyl ester (9CI) (CA INDEX NAME) |        |          |                 |          |

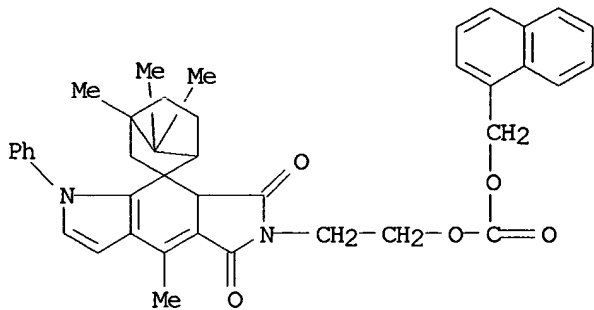


IT **123498-25-9P**

RL: PREP (Preparation)  
(prepn. of, as photochromic substance)

RN 123498-25-9 CAPLUS

CN Carbonic acid, 1-naphthalenylmethyl 2-(1,5,7,7a-tetrahydro-4,4',7',7'-tetramethyl-5,7-dioxo-1-phenylspiro[benzo[1,2-b:4,5-c']dipyrrole-8(6H),2'-bicyclo[2.2.1]heptan]-6-yl)ethyl ester (9CI) (CA INDEX NAME)



L12 ANSWER 30 OF 32 CAPLUS COPYRIGHT 2001 ACS  
 1989:173762 Document No. 110:173762 Preparation, testing, and formulation of

indol(in)e carboxylate-containing tripeptides as antihypertensives..  
 Sawayama, Tadahiro; Tsukamoto, Masatoshi; Sasagawa, Takashi; Nishimura, Kazuya; Hosoki, Kanoo; Takeyama, Kunihiko (Dainippon Pharmaceutical Co., Ltd., Japan). Eur. Pat. Appl. EP 244836 A2 19871111, 91 pp. DESIGNATED STATES: R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE.  
 (English). CODEN: EPXXDW. APPLICATION: EP 1987-106526 19870506.  
 PRIORITY: JP 1986-107394 19860509; JP 1986-156693 19860703; JP 1987-16361 19870126.

|    | PATENT NO.  | KIND | DATE     | APPLICATION NO. | DATE     |
|----|---|------|----------|-----------------|----------|
| PI | EP 244836   | A2   | 19871111 | EP 1987-106526  | 19870506 |
|    | EP 244836   | A3   | 19891123 |                 |          |
|    | EP 244836   | B1   | 19930818 |                 |          |
|    | R: AT, BE, CH, DE, ES, FR, GB, GR, IT, LI, LU, NL, SE |      |          |                 |          |
|    | AU 8772416  | A1   | 19871112 | AU 1987-72416   | 19870501 |
|    | AU 595309   | B2   | 19900329 |                 |          |
|    | US 4826814  | A    | 19890502 | US 1987-46189   | 19870505 |
|    | CA 1318461  | A1   | 19930525 | CA 1987-536368  | 19870505 |
|    | ZA 8703226  | A    | 19880427 | ZA 1987-3226    | 19870506 |
|    | AT 93237  | E    | 19930915 | AT 1987-106526  | 19870506 |
|    | ES 2058074  | T3   | 19941101 | ES 1987-106526  | 19870506 |
|    | DK 8702357  | A    | 19871110 | DK 1987-2357    | 19870508 |
|    | DK 171402   | B1   | 19961014 |                 |          |
|    | FI 8702041  | A    | 19871110 | FI 1987-2041    | 19870508 |
|    | FI 87794  | B    | 19921113 |                 |          |
|    | FI 87794  | C    | 19930225 |                 |          |
|    | DD 256329   | A5   | 19880504 | DD 1987-302570  | 19870508 |
|    | HU 45268  | A2   | 19880628 | HU 1987-2089    | 19870508 |
|    | HU 202884   | B    | 19910429 |                 |          |
|    | JP 63295597   | A2   | 19881201 | JP 1987-112831  | 19870508 |
|    | JP 05037998   | B4   | 19930607 |                 |          |
|    | SU 1743356  | A3   | 19920623 | SU 1987-4202607 | 19870508 |
|    | SK 278137   | B6   | 19960207 | SK 1987-3323    | 19870508 |
|    | CZ 280776   | B6   | 19960417 | CZ 1987-3323    | 19870508 |

IT 116587-40-7P

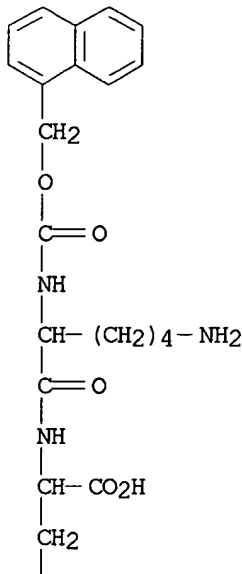
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)  
 (prepn. of, as antihypertensive)

RN 116587-40-7 CAPLUS

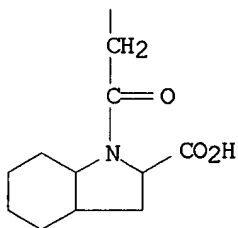
CN D-Norvaline, 5-(2-carboxyoctahydro-1H-indol-1-yl)-N-[N2-[(1-naphthalenylmethoxy)carbonyl]-L-lysyl]-5-oxo-, [2S-

(2.alpha.,3a.beta.,7a.beta.)]- (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A



L12 ANSWER 31 OF 32 CAPLUS COPYRIGHT 2001 ACS

1985:221199 Document No. 102:221199 Carboxyalkyl peptide derivatives.  
McCullagh, Keith; Wadsworth, Harry; Hann, Michael (Searle, G. D., and  
Co.,

USA). Eur. Pat. Appl. EP 126974 A1 19841205, 111 pp. DESIGNATED STATES:  
R: BE, CH, DE, FR, GB, IT, LI, NL, SE. (English). CODEN: EPXXDW.

APPLICATION: EP 1984-104614 19840425. PRIORITY: GB 1983-11286 19830426.

PATENT NO. KIND DATE APPLICATION NO. DATE

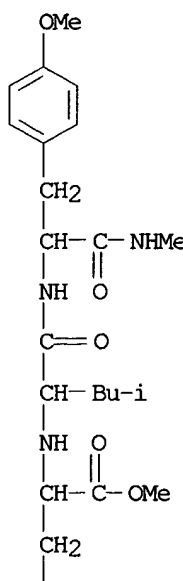
| PI                                    | EP 126974   | A1 | 19841205 | EP 1984-104614 | 19840425 |
|---------------------------------------|-------------|----|----------|----------------|----------|
|                                       | EP 126974   | B1 | 19880615 |                |          |
| R: BE, CH, DE, FR, GB, IT, LI, NL, SE |             |    |          |                |          |
|                                       | AU 8427222  | A1 | 19841122 | AU 1984-27222  | 19840424 |
|                                       | AU 575048   | B2 | 19880721 |                |          |
|                                       | ZA 8403056  | A  | 19850626 | ZA 1984-3056   | 19840425 |
|                                       | CA 1284850  | A1 | 19910611 | CA 1984-452746 | 19840425 |
|                                       | JP 59205350 | A2 | 19841120 | JP 1984-85091  | 19840426 |

JP 06045635 B4 19940615  
 JP 06316594 A2 19941115 JP 1993-256172 19931013  
 JP 2725690 B2 19980311  
 JP 08259593 A2 19961008 JP 1996-35137 19960222  
 JP 2706646 B2 19980128

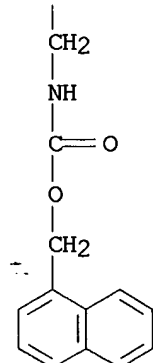
**IT 96134-98-4P 96134-99-5P**  
 RL: SPN (Synthetic preparation); PREP (Preparation)  
 (prepn. of)

**RN 96134-98-4 CAPLUS**  
**CN L-Tyrosinamide, N-[1-(methoxycarbonyl)-3-[(1-naphthalenylmethoxy)carbonyl]amino]propyl-L-leucyl-N,O-dimethyl-, (R)-(9CI) (CA INDEX NAME)**

PAGE 1-A



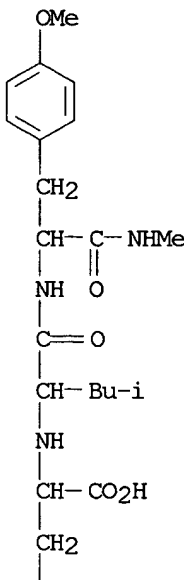
PAGE 2-A



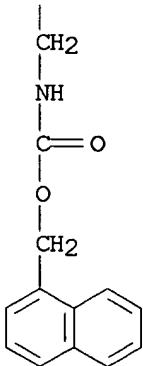
**RN 96134-99-5 CAPLUS**  
**CN L-Tyrosinamide,**  
**N-[1-carboxy-3-[(1-naphthalenylmethoxy)carbonyl]amino]pro**

pyl]-L-leucyl-N,O-dimethyl-, (R)- (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A



L12 ANSWER 32 OF 32 CAPLUS COPYRIGHT 2001 ACS

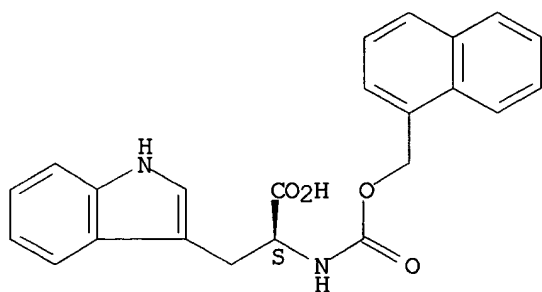
1973:72593 Document No. 78:72593 N-Acyl-L-, D-, and DL-tryptophan, their esters and amides for the treatment of gastric ulcers. Rovati, Luigi S.; Picciola, Giampaolo; Makovec, Francesco (Rotta Research Laboratorium). Ger. Offen. DE 2224130 19721130, 25 pp. (German). CODEN: GWXXBX.  
PRIORITY: IT 1971-68652 19710518.

| PATENT NO.    | KIND | DATE     | APPLICATION NO. | DATE     |
|---------------|------|----------|-----------------|----------|
| PI DE 2224130 | A    | 19721130 | DE 1972-2224130 | 19720517 |
| DE 2224130    | B2   | 19791108 |                 |          |
| DE 2224130    | C3   | 19800717 |                 |          |
| GB 1352472    | A    | 19740508 | GB 1972-11910   | 19720314 |

|             |    |          |                |          |
|-------------|----|----------|----------------|----------|
| ES 402678   | A1 | 19750401 | ES 1972-402678 | 19720512 |
| NL 7206680  | A  | 19721121 | NL 1972-6680   | 19720517 |
| NL 173167   | B  | 19830718 |                |          |
| NL 173167   | C  | 19831216 |                |          |
| FR 2138046  | A5 | 19721229 | FR 1972-17675  | 19720517 |
| FR 2138046  | B1 | 19750620 |                |          |
| JP 48048462 | A2 | 19730709 | JP 1972-48276  | 19720517 |
| JP 51039220 | B4 | 19761026 |                |          |
| US 4000297  | A  | 19761228 | US 1976-648359 | 19760112 |

IT **39545-08-9P**  
 RL: SPN (Synthetic preparation); PREP (Preparation)  
 (prepn. of)  
 RN 39545-08-9 CAPLUS  
 CN L-Tryptophan, N-[(1-naphthalenylmethoxy)carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



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ALL L# QUERIES AND ANSWER SETS ARE DELETED AT LOGOFF

LOGOFF? (Y)/N/HOLD:.

COST IN U.S. DOLLARS

FULL ESTIMATED COST

SINCE FILE  
ENTRY

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TOTAL  
SESSION  
310.08

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